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TITLE OF THE INVENTION

RECOMBINANT PLANT VIRAL NUCLEIC ACIDS

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CROSS-REFERENCE TO RELATED APPLICATIONS

The present application is a continuation-in-part of applications Serial No. 600,244, filed October 22, 1990, Serial No. 641,617, filed January 16, 1991, and Serial No. 737,899 filed July 26, 1991. Serial No. 600,244 is a continuation of application Serial No. 310,881, filed February 17, 1989, now abandoned, which is a continuation-in-part of applications Serial No. 160,766 and 160,771, both filed on February 26, 1988 and now abandoned. Serial No. 641,617 is a continuation of application Serial No. 347,637, filed May 5, 1989, now abandoned. Serial No. 737,899 is a continuation of application Serial No. 363,138, filed June 8, 1989, now abandoned, which is a continuation-in-part of application Serial No. 219,279, filed July 15, 1988 and now abandoned. The disclosures of all of the foregoing are incorporated by reference.

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BACKGROUND OF THE INVENTION

The present invention relates to plant viral vectors which are (a) self-replicating; (b) capable of systemic infection in a host; (c) contain, or are capable of containing, nucleic acid sequences foreign to the native virus, which are transcribed or expressed in the host plant; and (d) stable, especially for the transcription and expression of foreign nucleic acid sequences.

Viruses are a unique class of infectious agents whose distinctive features are their simple

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organization and their mechanism of replication. In fact, a complete viral particle, or virion, may be regarded mainly as a block of genetic material (either DNA or RNA) capable of autonomous replication, surrounded by a protein coat and sometimes by an additional membranous envelope such as in the case of alpha viruses. The coat protects the virus from the environment and serves as a vehicle for transmission from one host cell to another.

Unlike cells, viruses do not grow in size and then divide, because they contain within their coats few (or none) of the biosynthetic enzymes and other machinery required for their replication. Rather, viruses multiply in cells by the synthesis of their separate components, followed by assembly. Thus, the viral nucleic acid, after shedding its coat, comes into contact with the appropriate cell machinery where it specifies the synthesis of proteins required for viral reproduction. The viral nucleic acid is then itself replicated through the use of both viral and cellular enzymes. The components of the viral coat are formed and the nucleic acid and coat components are finally assembled. With some viruses, replication is initiated by enzymes present in virions.

A given plant virus may contain either DNA or RNA, which may be either single- or double-stranded. The portion of nucleic acid in a virion varies from about 1% to about 50%. The amount of genetic information per virion varies from about 3 kb to 300 kb per strand. The diversity of virus-specific proteins varies accordingly. One example of double-stranded DNA containing plant viruses includes, but is not limited to, caulimoviruses such as Cauliflower

mosaic virus (CaMV). Representative plant viruses which contain single-stranded DNA are Cassava latent virus, bean golden mosaic virus (BGMV), and Chloris striate mosaic virus. Rice dwarf virus and wound tumor virus are examples of double-stranded RNA plant viruses. Single-stranded RNA plant viruses include tobacco mosaic virus (TMV), turnip yellow mosaic virus (TYMV), rice necrosis virus (RNV) and brome mosaic virus (BMV). The RNA in single-stranded RNA viruses may be either a plus (+) or a minus (-) strand. For general information concerning plant viruses, see Grierson, D. et al. (1); Gluzman, Y. et al. (2).

One means for classifying plant viruses is based on the genome organization. Although many plant viruses have RNA genomes, organization of genetic information differs between groups. The genome of most monopartite plant RNA viruses is a single-stranded molecule of (+)- sense. There are at least 11 major groups of viruses with this type of genome. An example of this type of virus is TMV. At least six major groups of plant RNA viruses have a bipartite genome. In these, the genome usually consists of two distinct (+)- sense single-stranded RNA molecules encapsidated in separate particles. Both RNAs are required for infectivity. Cowpea mosaic virus (CPMW) is one example of a bipartite plant virus. A third major group, containing at least six major types of plant viruses, is tripartite, with three (+)- sense single-stranded RNA molecules. Each strand is separately encapsidated, and all three are required for infectivity. An example of a tripartite plant virus is alfalfa mosaic virus (AMV). Many plant viruses also have smaller subgenomic mRNAs that are synthesized to amplify a

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specific gene product. One group of plant viruses having a single-stranded DNA genome are the geminiviruses, such as Cassava latent virus (CLV) and maize streak virus (MSV). Several plant viruses have been cloned to study their nucleic acid, in anticipation of their use as plant transformation vectors. Examples of viruses cloned include BMV, Ahlgquist, P. and Janda, M. (3); TMV, Dawson W.O. et al. (4); CaMV, Lebeurier, G. et al. (5); and BGMV, Morinaga, T. et al. (6).

Techniques have been developed which are utilized to transform many species of organisms. Hosts which are capable of being transformed by these techniques include bacteria, yeast, fungus, animal cells and plant cells or tissue. Transformation is accomplished by using a vector which is self-replicating and which is compatible with the desired host. The vectors are generally based on either a plasmid or a virus. Foreign DNA is inserted into the vector, which is then used to transform the appropriate host. The transformed host is then identified by selection or screening. For further information concerning the transformation of these hosts, see Molecular Cloning (7) DNA Cloning (8); Grierson, D. et al. (1), and Methods in Enzymology, (9).

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Viruses that have been shown to be useful for the transformation of plant hosts include CaV, TMV and BV. Transformation of plants using plant viruses is described in US 4,855,237 (BGV), EP-A 67,553 (TMV), Japanese Published Application No. 63-14693 (TMV), EPA 194,809 (BV), EPA 278,667 (BV), Brisson, N. et al. (10) (CaV), and Guzman et al. (2). Pseudovirus particles for use in expressing foreign

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DNA in many hosts, including plants, is described in WO 87/06261.

When the virus is a DNA virus, the constructions can be made to the virus itself. Alternatively, the virus can first be cloned into a bacterial plasmid for ease of constructing the desired viral vector with the foreign DNA. The virus can then be excised from the plasmid. If the virus is a DNA virus, a bacterial origin of replication can be attached to the viral DNA, which is then replicated by the bacteria. Transcription and translation of this DNA will produce the coat protein which will encapsidate the viral DNA. If the virus is an RNA virus, the virus is generally cloned as a cDNA and inserted into a plasmid. The plasmid is then used to make all of the constructions. The RNA virus is then produced by transcribing the viral sequence of the plasmid and translation of the viral genes to produce the coat protein(s) which encapsidate the viral RNA.

Construction of plant RNA viruses for the introduction and expression of non-viral foreign genes in plants is demonstrated by the above references as well as by Dawson, W.O. et al. (11); Takamatsu, N. et al. (12); French, R. et al. (13); and Takamatsu, N. et al. (14). However, none of these viral vectors have been capable of systemic spread in the plant and expression of the non-viral foreign genes in the majority of the plant cells in the whole plant. Another disadvantage of many of the prior art viral vectors is that they are not stable for the maintenance of non-viral foreign genes. See, for example, Dawson, W.O. et al. (11),. Thus, despite all of this activity to develop plant viral vectors and viruses, a need still exists for a stable recombinant plant virus capable of systemic infection

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in the host plant and stable expression of the foreign DNA.

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SUMMARY OF THE INVENTION

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The present invention is directed to recombinant plant viral nucleic acids and recombinant viruses which are stable for maintenance and transcription or expression of non-native (foreign) nucleic acid sequences and which are capable of systemically transcribing or expressing such foreign sequences in the host plant. More specifically, recombinant plant viral nucleic acids according to the present invention comprise a native plant viral subgenomic promoter, at least one non-native plant viral subgenomic promoter, a plant viral coat protein coding sequence, and optionally, at least one non-native, nucleic acid sequence.

In one embodiment, a plant viral nucleic acid is provided in which the native coat protein coding sequence has been deleted from a viral nucleic acid, a non-native plant viral coat protein coding sequence and a non-native promoter, preferably the subgenomic promoter of the non-native coat protein coding sequence, capable of expression in the plant host, packaging of the recombinant plant viral nucleic acid, and ensuring a systemic infection of the host by the recombinant plant viral nucleic acid, has been inserted.

The recombinant plant viral nucleic acid may contain one or more additional non-native subgenomic promoters. Each non-native subgenomic promoter is capable of transcribing or expressing adjacent genes or nucleic acid sequences in the plant host and

incapable of recombination with each other and with native subgenomic promoters.

Non-native (foreign) nucleic acid sequences may be inserted adjacent the native plant viral subgenomic promoter or the native and a non-native plant viral subgenomic promoters if more than one nucleic acid sequence is included. The non-native nucleic acid sequences are transcribed or expressed in the host plant under control of the subgenomic promoter to produce the desired products.

In a second embodiment, a recombinant plant viral nucleic acid is provided as in the first embodiment except that the native coat protein coding sequence is placed adjacent one of the non-native coat protein subgenomic promoters instead of a non-native coat protein coding sequence.

In a third embodiment, a recombinant plant viral nucleic acid is provided in which the native coat protein gene is adjacent its subgenomic promoter and one or more non-native subgenomic promoters have been inserted into the viral nucleic acid. The inserted non-native subgenomic promoters are capable of transcribing or expressing adjacent genes in a plant host and are incapable of recombination with each other and with native subgenomic promoters. Non-native nucleic acid sequences may be inserted adjacent the non-native subgenomic plant viral promoters such that said sequences are transcribed or expressed in the host plant under control of the subgenomic promoters to produce the desired product.

In a fourth embodiment, a recombinant plant viral nucleic acid is provided as in the third embodiment except that the native coat protein coding sequence is replaced by a non-native coat protein coding sequence.

The viral vectors are encapsidated by the coat proteins encoded by the recombinant plant viral nucleic acid to produce a recombinant plant virus. The recombinant plant viral nucleic acid or recombinant plant virus is used to infect appropriate host plants. The recombinant plant viral nucleic acid is capable of replication in the host, systemic spread in the host, and transcription or expression of foreign gene(s) in the host to produce the desired product. Such products include therapeutic and other useful polypeptides or proteins such as, but not limited to, enzymes, complex biomolecules, ribozymes, or polypeptide or protein products resulting from anti-sense RNA expression.

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BRIEF DESCRIPTION OF THE FIGURES

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Figure 1 illustrates several vectors prepared in accordance with the present invention and restriction sites. U1 is the native plant viral nucleic acid, O is a non-native plant viral nucleic acid, and the hatched area is a non-native plant viral subgenomic promoter. The restriction sites are: X-XbaI, N-NsiI, K-KpnI, S-SplI, B-BamHI, Nc-NcoI, P-PstI. The hatched box (e.g., in TB2) represents the promoter of TMV-O, i.e., 203 bp upstream of the coat protein initiation codon, and the stippled box represents a phage promoter. The open boxes represent open reading frames, and the solid boxes represent cloning vector sequences. The vectors are as follows: A) and B) pTKU1, C) pTMVS3-28, D) pTB2, E) pTBN62 and F) pTBU5.

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Figure 2 is an autoradiograph of a Western analysis of the production of α -trichosanthin in N. benthamiana infected in accordance with the present

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invention. Lane a is molecular size markers, lanes b and c are extracts from yeast engineered to produce α-trichosanthin and lane d is a extract from N. benthamiana.

Figures 3^A and 3^B

Figures 3^A and 3^B illustrate the α-trichosanthin expression vector, pBGC152. This plasmid contains the TMV-U1 126-, 183-, and 30-kDa open reading frames (ORFs), the ORSV coat protein gene (Ocp), the SP6 promoter, the α-trichosanthin gene, and part of the pBR322 plasmid. The TAA stop codon in the 30K ORF is underlined and a bar (—) divides the putative signal peptide from the mature peptide. The TMV-U1 subgenomic promoter located within the minus strand of the 30K ORF controls the expression of α-trichosanthin. The putative transcription start point (tsp) of the subgenomic RNA is indicated with a period (.).

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Figure 4 illustrates an electron micrograph of virions from systemically infected leaves of N. benthamiana transfected with in vivo pBGC152 transcripts. The length of the black bar located in the bottom left corner of the micrograph represents approximately 140 nm.

Figure 5a is a protein analysis of a transfected N. benthamiana plant two weeks after inoculation. a, Western blot analysis. Lane 1: 200 ng of GLQ223; 2: 50 ng of GLQ223; 3: 7 μg of total soluble protein from N. benthamiana infected with pBGC152 transcripts; 4: peak fraction from alkyl superose FPLC chromatography; 5: 7 μg of total soluble protein from noninfected N. benthamiana; 6: 7 μg of total soluble protein from noninfected N. benthamiana and 100 ng of GLQ223.

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60 Figure 5b is a purification profile of recombinant α -trichosanthin. The samples from various stages during purification were analyzed by 12.5% SDS-polyacrylamide gel electrophoresis.

I Lane 1: Amersham prestained high-range molecular weight standards; 2: purified GLQ223; 3: total soluble protein from N. benthamiana infected with pBGC152 transcripts; 4: peak fraction from S-sepharose chromatography; 5: peak fraction from alkyl superose FPLC chromatography.

Figure 6 illustrates the inhibition of protein synthesis in a cell-free rabbit reticulocyte translation assay. Dosage required for 50%

H₆₀ inhibition (ID₅₀). Purified α -trichosanthin from I N. benthamiana infected with BGC 152 transcripts (blackened circles and triangles, repetition 1 and 2), GLQ233 (blackened square), and cycloheximide (open circle) were analyzed in varying concentrations for their ability to inhibit protein synthesis in vitro.

Figure 7 illustrates the construction of the pBGC152 plasmid.

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DETAILED DESCRIPTION OF THE INVENTION

The present invention is directed to recombinant plant viral nucleic acids and recombinant viruses which are stable for maintenance and transcription or expression of non-native (foreign) nucleic acid sequences and which are capable of systemically transcribing or expressing such foreign sequences in the host plant. More specifically, recombinant plant viral nucleic acids according to the present invention comprise a native plant viral subgenomic promoter, at least one non-native plant viral subgenomic promoter, a plant viral coat protein

coding sequence, and optionally, at least one non-native, nucleic acid sequence.

In one embodiment, a plant viral nucleic acid is provided in which the native coat protein coding sequence has been deleted from a viral nucleic acid, a non-native plant viral coat protein coding sequence and a non-native promoter, preferably the subgenomic promoter of the non-native coat protein coding sequence, capable of expression in the plant host, packaging of the recombinant plant viral nucleic acid, and ensuring a systemic infection of the host by the recombinant plant viral nucleic acid, has been inserted. Alternatively, the coat protein gene may be inactivated by insertion of the non-native nucleic acid sequence within it, such that a fusion protein is produced. The recombinant plant viral nucleic acid may contain one or more additional non-native subgenomic promoters. Each non-native subgenomic promoter is capable of transcribing or expressing adjacent genes or nucleic acid sequences in the plant host and incapable of recombination with each other and with native subgenomic promoters. Non-native (foreign) nucleic acid sequences may be inserted adjacent the native plant viral subgenomic promoter or the native and a non-native plant viral subgenomic promoters if more than one nucleic acid sequence is included. The non-native nucleic acid sequences are transcribed or expressed in the host plant under control of the subgenomic promoter to produce the desired products.

In a second embodiment, a recombinant plant viral nucleic acid is provided as in the first embodiment except that the native coat protein coding sequence is placed adjacent one of the non-native

coat protein subgenomic promoters instead of a non-native coat protein coding sequence.

In a third embodiment, a recombinant plant viral nucleic acid is provided in which the native coat protein gene is adjacent its subgenomic promoter and one or more non-native subgenomic promoters have been inserted into the viral nucleic acid. The inserted non-native subgenomic promoters are capable of transcribing or expressing adjacent genes in a plant host and are incapable of recombination with each other and with native subgenomic promoters. Non-native nucleic acid sequences may be inserted adjacent the non-native subgenomic plant viral promoters such that said sequences are transcribed or expressed in the host plant under control of the subgenomic promoters to produce the desired product.

In a fourth embodiment, a recombinant plant viral nucleic acid is provided as in the third embodiment except that the native coat protein coding sequence is replaced by a non-native coat protein coding sequence.

The viral vectors are encapsidated by the coat proteins encoded by the recombinant plant viral nucleic acid to produce a recombinant plant virus. The recombinant plant viral nucleic acid or recombinant plant virus is used to infect appropriate host plants. The recombinant plant viral nucleic acid is capable of replication in the host, systemic spread in the host, and transcription or expression of foreign gene(s) in the host to produce the desired product.

In order to provide a clear and consistent understanding of the specification and the claims, including the scope given herein to such terms, the following definitions are provided:

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Adjacent: A position in a nucleotide sequence immediately 5' or 3' to a defined sequence.

Anti-Sense Mechanism: A type of gene regulation based on controlling the rate of translation of mRNA to protein due to the presence in a cell of an RNA molecule complementary to at least a portion of the mRNA being translated.

Cell Culture: A proliferating mass of cells which may be in either an undifferentiated or differentiated state.

Chimeric Sequence or Gene: A nucleotide sequence derived from at least two heterologous parts. The sequence may comprise DNA or RNA.

Coding Sequence: A deoxyribonucleotide sequence which, when transcribed and translated, results in the formation of a cellular polypeptide or a ribonucleotide sequence which, when translated, results in the formation of a cellular polypeptide.

Compatible: The capability of operating with other components of a system. A vector or plant viral nucleic acid which is compatible with a host is one which is capable of replicating in that host. A coat protein which is compatible with a viral nucleotide sequence is one capable of encapsidating that viral sequence.

Gene: A discrete nucleic acid sequence responsible for a discrete cellular product.

Host: A cell, tissue or organism capable of replicating a vector or plant viral nucleic acid and which is capable of being infected by a virus containing the viral vector or plant viral nucleic acid. This term is intended to include prokaryotic and eukaryotic cells, organs, tissues or organisms, where appropriate.

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Infection: The ability of a virus to transfer its nucleic acid to a host or introduce viral nucleic acid into a host, wherein the viral nucleic acid is replicated, viral proteins are synthesized, and new viral particles assembled. In this context, the terms "transmissible" and "infective" are used interchangeably herein.

Non-Native: Any RNA sequence that promotes production of subgenomic mRNA including, but not limited to, 1) plant viral promoters such as ORSV and bromo mosaic virus, 2) viral promoters from other organisms such as human sindbis viral promoter, and 3) synthetic promoters.

Phenotypic Trait: An observable property resulting from the expression of a gene.

Plant Cell: The structural and physiological unit of plants, consisting of a protoplast and the cell wall.

Plant Organ: A distinct and visibly differentiated part of a plant, such as root, stem, leaf or embryo.

Plant Tissue: Any tissue of a plant in planta or in culture. This term is intended to include a whole plant, plant cell, plant organ, protoplast, cell culture, or any group of plant cells organized into a structural and functional unit.

Production Cell: A cell, tissue or organism capable of replicating a vector or a viral vector, but which is not necessarily a host to the virus. This term is intended to include prokaryotic and eukaryotic cells, organs, tissues or organisms, such as bacteria, yeast, fungus and plant tissue.

Promoter: The 5'-flanking, non-coding sequence adjacent a coding sequence which is involved in the initiation of transcription of the coding sequence.

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Protoplast: An isolated plant cell without cell walls, having the potency for regeneration into cell culture or a whole plant.

Recombinant Plant Viral Nucleic Acid: Plant viral nucleic acid which has been modified to contain nonnative nucleic acid sequences.

Recombinant Plant Virus: A plant virus containing the recombinant plant viral nucleic acid.

Subgenomic Promoter: A promoter of a subgenomic mRNA of a viral nucleic acid.

Substantial Sequence Homology: Denotes nucleotide sequences that are substantially functionally equivalent to one another. Nucleotide differences between such sequences having substantial sequence homology will be de minimus in affecting function of the gene products or an RNA coded for by such sequence.

Transcription: Production of an RNA molecule by RNA polymerase as a complementary copy of a DNA sequence.

Vector: A self-replicating DNA molecule which transfers a DNA segment between cells.

Virus: An infectious agent composed of a nucleic acid encapsidated in a protein. A virus may be a mono-, di-, tri- or multi-partite virus, as described above.

The present invention provides for the infection of a plant host by a recombinant plant virus containing recombinant plant viral nucleic acid or by the recombinant plant viral nucleic acid which contains one or more non-native nucleic acid sequences which are transcribed or expressed in the infected tissues of the plant host. The product of the coding sequences may be recovered from the plant

or cause a phenotypic trait, such as male sterility, in the plant.

The present invention has a number of advantages, one of which is that the transformation and regeneration of target organisms is unnecessary. Another advantage is that it is unnecessary to develop vectors which integrate a desired coding sequence in the genome of the target organism. Existing organisms can be altered with a new coding sequence without the need of going through a germ cell. The present invention also gives the option of applying the coding sequence to the desired organism, tissue, organ or cell. Recombinant plant viral nucleic acid is also stable for the foreign coding sequences, and the recombinant plant virus or recombinant plant viral nucleic acid is capable of systemic infection in the plant host.

Chimeric genes and vectors and recombinant plant viral nucleic acids according to this invention are constructed using techniques well known in the art.

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Suitable techniques have been described in Molecular Cloning (7); Methods in Enzymol. (9); and DNA Cloning (8). Medium compositions have been described in Miller, J.H. (15), as well as the references previously identified. DNA manipulations and enzyme treatments are carried out in accordance with manufacturers' recommended procedures.

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An important feature of the present invention is the preparation of recombinant plant viral nucleic acids (RPVNA) which are capable of replication and systemic spread in a compatible plant host, and which contain one or more non-native subgenomic promoters which are capable of transcribing or expressing adjacent nucleic acid sequences in the plant host. The RPVNA may be further modified to delete all or

part of the native coat protein coding sequence and to contain a non-native coat protein coding sequence under control of the native or one of the non-native subgenomic promoters, or put the native coat protein coding sequence under the control of a non-native plant viral subgenomic promoter. The RPVNA have substantial sequence homology to plant viral nucleotide sequences. A partial listing of suitable viruses has been described above. The nucleotide sequence may be an RNA, DNA, cDNA or chemically synthesized RNA or DNA.

The first step in achieving any of the features of the invention is to modify the nucleotide sequences of the plant viral nucleotide sequence by known conventional techniques such that one or more non-native subgenomic promoters are inserted into the plant viral nucleic acid without destroying the biological function of the plant viral nucleic acid. The subgenomic promoters are capable of transcribing or expressing adjacent nucleic acid sequences in a plant host infected by the recombinant plant viral nucleic acid or recombinant plant virus. The native coat protein coding sequence may be deleted in two embodiments, placed under the control of a non-native subgenomic promoter in a second embodiment, or retained in a further embodiment. If it is deleted or otherwise inactivated, a non-native coat protein gene is inserted under control of one of the non-native subgenomic promoters, or optionally under control of the native coat protein gene subgenomic promoter. The non-native coat protein is capable of encapsidating the recombinant plant viral nucleic acid to produce a recombinant plant virus. Thus, the recombinant plant viral nucleic acid contains a coat protein coding sequence, which may be native or a

nonnative coat protein coding sequence, under control of one of the native or non-native subgenomic promoters. The coat protein is involved in the systemic infection of the plant host.

Some of the viruses which meet this requirement, and are therefore suitable, include viruses from the tobacco mosaic virus group such as Tobacco Mosaic virus (TMV), Cowpea Mosaic virus (CMV), Alfalfa Mosaic virus (AMV), Cucumber Green Mottle Mosaic virus watermelon strain (CGMMV-W) and Oat Mosaic virus (OMV) and viruses from the brome mosaic virus group such as Brome Mosaic virus (MBV), broad bean mottle virus and cowpea chlorotic mottle virus. Additional suitable viruses include Rice Necrosis virus (RNV), and geminiviruses such as tomato golden mosaic virus (TGMV), Cassava latent virus (CLV) and maize streak virus (MSV). Each of these groups of suitable viruses is characterized below.

Tobacco Mosaic Virus Group

Tobacco Mosaic virus (TMV) is a member of the Tobamoviruses. The TMV virion is a tubular filament, and comprises coat protein sub-units arranged in a single right-handed helix with the single-stranded RNA intercalated between the turns of the helix. TMV infects tobacco as well as other plants. TMV is transmitted mechanically and may remain infective for a year or more in soil or dried leaf tissue.

The TMV virions may be inactivated by subjection to an environment with a pH of less than 3 or greater than 8, or by formaldehyde or iodine. Preparations of TMV may be obtained from plant tissues by $(\text{NH}_4)_2\text{SO}_4$ precipitation, followed by differential centrifugation.

40 The TMV single-stranded RNA genome is about 6400 nucleotides long, and is capped at the 5' end but not polyadenylated. The genomic RNA can serve as mRNA for a protein of a molecular weight of about 130,000 (130K) and another produced by read-through of molecular weight about 180,000 (180K). However, it cannot function as a messenger for the synthesis of coat protein. Other genes are expressed during infection by the formation of monocistronic, 40 3'-coterminal sub-genomic mRNAs, including one (LMC) H encoding the 17.5K coat protein and another (I_2) encoding a 30K protein. The 30K protein has been detected in infected protoplasts (16), and it is involved in the cell-to-cell transport of the virus in an infected plant (17). The functions of the two large proteins are unknown.

H Several double-stranded RNA molecules, including double-stranded RNAs corresponding to the genomic, I_2 and LMC RNAs, have been detected in plant tissues infected with TMV. These RNA molecules are presumably intermediates in genome replication and/or mRNA synthesis processes which appear to occur by different mechanisms.

TMV assembly apparently occurs in plant cell cytoplasm, although it has been suggested that some TMV assembly may occur in chloroplasts since transcripts of ctDNA have been detected in purified TMV virions. Initiation of TMV assembly occurs by interaction between ring-shaped aggregates ("discs") of coat protein (each disc consisting of two layers of 17 subunits) and a unique internal nucleation site in the RNA; a hairpin region about 900 nucleotides from the 3' end in the common strain of TMV. Any RNA, including subgenomic RNAs containing this site, may be packaged into virions. The discs apparently

assume a helical form on interaction with the RNA, and assembly (elongation) then proceeds in both directions (but much more rapidly in the 3'- to 5'- O direction from the nucleation site).

Another member of the Tobamoviruses, the Cucumber green mottle mosaic virus watermelon strain (CGMMV-W) is related to the cucumber virus. Noru, Y. et al. (18). The coat protein of CGMMV-W interacts with RNA of both TMV and CGMMV to assemble viral particles in vitro. Kurisu et al. (19).

Several strains of the tobamovirus group are divided into two subgroups, on the basis of the location of the assembly of origin. Fukuda, M. et al. (20). Subgroup I, which includes the vulgare, OM, and tomato strain, has an origin of assembly about 800-1000 nucleotides from the 3' end of the RNA genome, and outside the coat protein cistron.

Lebeurier, G. et al. (21); and Fukuda, M. et al. (22). Subgroup II, which includes CGMMV-W and cornpea strain (Cc) has an origin of assembly about 300-500 nucleotides from the 3' end of the RNA genome and within the coat-protein cistron. Fukuda, M. et al. (22). The coat protein cistron of CGMMV-W is located at nucleotides 176-661 from the 3' end. The 3' noncoding region is 175 nucleotides long. The origin of assembly is positioned within the coat protein cistron. Meshi, T. et al. (23).

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Brome Mosaic Virus Group

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Brome mosaic virus (BV) is a member of a group of tripartite, single-stranded, RNA-containing plant viruses commonly referred to as the bromoviruses. Each member of the bromoviruses infects a narrow range of plants. Mechanical transmission of

bromoviruses occurs readily, and some members are transmitted by beetles. In addition to BV, other bromoviruses include broad bean mottle virus and cowpea chlorotic mottle virus.

Typically, a bromovirus virion is icosahedral, with a diameter of about 26 nm, containing a single species of coat protein. The bromovirus genome has three molecules of linear, positive-sense, single-stranded RNA, and the coat protein mRNA is also encapsidated. The RNAs each have a capped 5' end, and a tRNA-like structure (which accepts tyrosine) at the 3' end. Virus assembly occurs in the cytoplasm. The complete nucleotide sequence of BMV has been identified and characterized as described by Alquist et al. (24).

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Rice Necrosis Virus

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Rice Necrosis virus is a member of the Potato Virus Y Group or Potyviruses. The Rice Necrosis virion is a flexuous filament comprising one type of coat protein (molecular weight about 32,000 to about 36,000) and one molecule of linear positive-sense single-stranded RNA. The Rice Necrosis virus is transmitted by Polymvxa araminis (a eukaryotic intracellular parasite found in plants, algae and fungi).

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Geminiviruses

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Geminiviruses are a group of small, single-stranded DNA-containing plant viruses with virions of unique morphology. Each virion consists of a pair of isometric particles (incomplete icosahedra), composed of a single type of protein (with a molecular weight

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of about $2.7\text{--}3.4 \times 10^4$). Each geminivirus virion contains one molecule of circular, positive-sense, single-stranded DNA. In some geminiviruses (i.e., Cassava latent virus and bean golden mosaic virus) the genome appears to be bipartite, containing two single-stranded DNA molecules.

The nucleic acid of any suitable plant virus can be utilized to prepare the recombinant plant viral nucleic acid of the present invention. The nucleotide sequence of the plant virus is modified, using conventional techniques, by the insertion of one or more subgenomic promoters into the plant viral nucleic acid. The subgenomic promoters are capable of functioning in the specific host plant. For example, if the host is tobacco, TMV will be utilized. The inserted subgenomic promoters must be compatible with the TMV nucleic acid and capable of directing transcription or expression of adjacent nucleic acid sequences in tobacco.

The native coat protein gene could also be retained and a non-native nucleic acid sequence inserted within it to create a fusion protein as discussed below. In this example, a non-native coat protein gene is also utilized.

The native or non-native coat protein gene is utilized in the recombinant plant viral nucleic acid. Whichever gene is utilized may be positioned adjacent its natural subgenomic promoter or adjacent one of the other available subgenomic promoters. The non-native coat protein, as is the case for the native coat protein, is capable of encapsidating the recombinant plant viral nucleic acid and providing for systemic spread of the recombinant plant viral nucleic acid in the host plant. The coat protein is selected to provide a systemic infection in the plant

I host of interest. For example, the TMV-O coat protein provides systemic infection in N. benthamiana, whereas TMV-U1 coat protein provides systemic infection in N. tabacum.

The recombinant plant viral nucleic acid is prepared by cloning viral nucleic acid in an appropriate production cell. If the viral nucleic acid is DNA, it can be cloned directly into a suitable vector using conventional techniques. One technique is to attach an origin of replication to the viral DNA which is compatible with the production cell. If the viral nucleic acid is RNA, a full-length DNA copy of the viral genome is first prepared by well-known procedures. For example, the viral RNA is transcribed into DNA using reverse transcriptase to produce subgenomic DNA pieces, and a double-stranded DNA made using DNA polymerases. The DNA is then cloned into appropriate vectors and cloned into a production cell. The DNA pieces are mapped and combined in proper sequence to produce a full-length DNA copy of the viral RNA genome, if necessary. DNA sequences for the subgenomic promoters, with or without a coat protein gene, are then inserted into the nucleic acid at non-essential sites, according to the particular embodiment of the invention utilized. Non-essential sites are those that do not affect the biological properties of the plant viral nucleic acid. Since the RNA genome is the infective agent, the cDNA is positioned adjacent a suitable promoter so that the RNA is produced in the production cell. The RNA is capped using conventional techniques, if the capped RNA is the infective agent.

A second feature of the present invention is a recombinant plant viral nucleic acid which further comprises one or more non-native nucleic acid

sequences capable of being transcribed in the plant host. The non-native nucleic acid sequence is placed adjacent one or the non-native viral subgenomic promoters and/or the native coat protein gene promoter depending on the particular embodiment used. The non-native nucleic acid is inserted by conventional techniques, or the non-native nucleic acid sequence can be inserted into or adjacent the native coat protein coding sequence such that a fusion protein is produced. The non-native nucleic acid sequence which is transcribed may be transcribed as an RNA which is capable of regulating the expression of a phenotypic trait by an anti-sense mechanism. Alternatively, the non-native nucleic acid sequence in the recombinant plant viral nucleic acid may be transcribed and translated in the plant host, to produce a phenotypic trait. The non-native nucleic acid sequence(s) may also code for the expression of more than one phenotypic trait. The recombinant plant viral nucleic acid containing the non-native nucleic acid sequence is constructed using conventional techniques such that non-native nucleic acid sequence(s) are in proper orientation to whichever viral subgenomic promoter is utilized.

Useful phenotypic traits in plant cells include, but are not limited to, improved tolerance to herbicides, improved tolerance to extremes of heat or cold, drought, salinity or osmotic stress; improved resistance to pests (insects, nematodes or arachnids) or diseases (fungal, bacterial or viral) production of enzymes or secondary metabolites; male or female sterility; dwarfness; early maturity; improved yield, vigor, heterosis, nutritional qualities, flavor or processing properties, and the like. Other examples include the production of important proteins or other

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products for commercial use, such as lipase, melanin, pigments, antibodies, hormones, pharmaceuticals, antibiotics and the like. Another useful phenotypic trait is the production of degradative or inhibitory enzymes, such as are utilized to prevent or inhibit root development in malting barley. The phenotypic trait may also be a secondary metabolite whose production is desired in a bioreactor.

A double-stranded DNA of the recombinant plant viral nucleic acid or a complementary copy of the recombinant plant viral nucleic acid is cloned into a production cell. If the viral nucleic acid is an RNA molecule, the nucleic acid (cDNA) is first attached to a promoter which is compatible with the production cell. The RPVNA can then be cloned into any suitable vector which is compatible with the production cell. In this manner, only RNA copies of the chimeric nucleotide sequence are produced in the production cell. For example, if the production cell is E. coli, the lac promoter can be utilized. If the production cell is a plant cell, the CaMV promoter can be used. The production cell can be a eukaryotic cell such as yeast, plant or animal, if viral RNA must be capped for biological activity. Alternatively, the RPVNA is inserted in a vector adjacent a promoter which is compatible with the production cell. If the viral nucleic acid is a DNA molecule, it can be cloned directly into a production cell by attaching it to an origin of replication which is compatible with the production cell. In this manner, DNA copies of the chimeric nucleotide sequence are produced in the production cell.

A promoter is a DNA sequence that directs RNA polymerase to bind to DNA and to initiate RNA synthesis. There are strong promoters and weak

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80 promoters. Among the strong promoters are lacuv5, trp, tac, trp-lacuv5, λ p1, ompF, and bla. A useful promoter for expressing foreign genes in E. coli is one which is both strong and regulated. The λ p1 promoter of bacteriophage λ is a strong, well-regulated promoter. Hedgpeth, J.M. et al. (25); Bernard, H.M. et al. (26); Remaut, E.P. et al. (27).

80 L A gene encoding a temperature-sensitive λ repressor such as λ cIIts 857 may be included in the cloning vector. Bernard et al. (26). At low temperature (31°C), the p₁ promoter is maintained in a repressed state by the cI-gene product. Raising the temperature destroys the activity of the repressor.

H
I The p₁ promoter then directs the synthesis of large quantities of mRNA. In this way, E. coli production cells may grow to the desired concentration before producing the products encoded within the vectors. Similarly, a temperature-sensitive promoter may be activated at the desired time by adjusting the temperature of the culture.

62 It may be advantageous to assemble a plasmid that can conditionally attain very high copy numbers. For example, the pAS2 plasmid containing a lac or tac promoter will achieve very high copy numbers at 42°C . The lac repressor, present in the pAS2 plasmid, is then inactivated by isopropyl- β -D-thiogalactoside to allow synthesis of mRNA.

A further alternative when creating the RPVNA is to prepare more than one nucleic acid (i.e., to prepare the nucleic acids necessary for a multipartite viral vector construct). In this case, each nucleic acid would require its own origin of assembly. Each nucleic acid could be prepared to contain a subgenomic promoter and a non-native nucleic acid.

Alternatively, the insertion of a non-native nucleic acid into the nucleic acid of a monopartite virus may result in the creation of two nucleic acids (i.e., the nucleic acid necessary for the creation of a bipartite viral vector). This would be advantageous when it is desirable to keep the replication and transcription or expression of the non-native nucleic acid separate from the replication and translation of some of the coding sequences of the native nucleic acid. Each nucleic acid would have to have its own origin of assembly.

A third feature of the present invention is a virus or viral particle. The virus comprises a RPVNA as described above which has been encapsidated. The resulting product is then capable of infecting an appropriate plant host. The RPVNA sequence is transcribed and/or translated within the plant host to produce the desired product.

In one embodiment of the present invention, the recombinant plant viral nucleic acid is encapsidated by a heterologous capsid. Most commonly, this embodiment will make use of a rod-shaped capsid because of its ability to encapsidate a longer RPVNA than the more geometrically constrained icosahedral capsid or spherical capsid. The use of a rod-shaped capsid permits incorporation of a larger non-native nucleic acid to form the RPVNA. Such a rod-shaped capsid is most advantageous when more than one non-native nucleic acid is present in the RPVNA.

Another feature of the invention is a vector containing the RPVNA as described above. The RPVNA is adjacent a nucleotide sequence selected from the group consisting of a production cell promoter or an origin of replication compatible with the production cell. The vector is utilized to transform a

production cell which will then produce the RPVNA in quantity. The production cell may be any cell which is compatible with the vector, and may be prokaryotic or eukaryotic. However, if the viral RNA (RPVNA) must be capped in order to be active, the production cell must be capable of capping the viral RNA, such as a eukaryotic production cell.

A further feature of the present invention is a host which has been infected by the recombinant plant virus or viral nucleic acid. After introduction into a host, the host contains the RPVNA which is capable of self-replication, encapsidation and systemic spread. The host can be infected with the recombinant plant virus by conventional techniques. Suitable techniques include, but are not limited to, leaf abrasion, abrasion in solution, high velocity water spray and other injury of a host as well as imbibing host seeds with water containing the recombinant plant virus. More specifically, suitable techniques include:

(a) Hand Inoculations. Hand inoculations of the encapsidated vector are performed using a neutral pH, low molarity phosphate buffer, with the addition of celite or carborundum (usually about 1%). One to four drops of the preparation is put onto the upper surface of a leaf and gently rubbed.

(b) Mechanized Inoculations of Plant Beds.

H Plant bed inoculations are performed by spraying (CO₂-propelled) the vector solution into a tractor-driven mower while cutting the leaves.

Alternatively, the plant bed is mowed and the vector solution sprayed immediately onto the cut leaves.

(c) High Pressure Spray of Single Leaves.

Single plant inoculations can also be performed by spraying the leaves with a narrow, directed spray (50

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psi, 6-12 inches from the leaf) containing approximately 1% carborundum in the buffered vector solution.

An alternative method for introducing a RPVNA into a plant host is a technique known as agroinfection or Agrobacterium-mediated transformation (sometimes called Agro-infection) as described by Grimsley, N. et al. (28). This technique makes use of a common feature of Agrobacterium which colonizes plants by transferring a portion of their DNA (the T-DNA) into a host cell, where it becomes integrated into nuclear DNA. The T-DNA is defined by border sequences which are 25 base pairs long, and any DNA between these border sequences is transferred to the plant cells as well. The insertion of a RPVNA between the T-DNA border sequences results in transfer of the RPVNA to the plant cells, where the RPVNA is replicated, and then spreads systemically through the plant. Agro-infection has been accomplished with potato spindle tuber viroid (PSTV) (Gardner, R.C. et al. (29)); CaV (Grimsley, N. et al. (30)); MSV (Grimsley, N. et al. (28), supra) and Lazarowitz, S.C. (31)), digitaria streak virus (Donson, J. et al. (32)), wheat dwarf virus (Hayes, R.J. et al. (33)) and tomato golden mosaic virus (TGMV) (Elmer, J.S. et al. (34) and Gardiner, W.E. et al. (35)). Therefore, agro-infection of a susceptible plant could be accomplished with a virion containing a RPVNA based on the nucleotide sequence of any of the above viruses.

A still further feature of the invention is a process for the production of a specified polypeptide or protein product such as, but not limited to, enzymes, complex biomolecules, a ribozyme, or

polypeptide or protein products resulting from anti-sense RNA. Such products include, but not limited to: IL-1, IL-2, IL-3, ... IL-12, etc.; EPO; CSF including G-CSF, GM-CSF, hPG-CSF, M-CSF, etc; Factor VIII; Factor IX; tPA; hGH; receptors and receptor antagonists; antibodies; neuro-polypeptides; melanin; insulin; vaccines and the like. The non-native nucleic acid of the RPVNA comprises the transcribable sequence which leads to the production of the desired product. This process involves the infection of the appropriate plant host with a recombinant virus or recombinant plant viral nucleic acid such as those described above, the growth of the infected host to produce the desired product, and the isolation of the desired product, if necessary. The growth of the infected host is in accordance with conventional techniques, as is the isolation of the resultant product.

For example, a coding sequence for a protein such as neomycin phosphotransferase (NPTII) ⁶⁰ ⁶² α -trichosanthin, rice α -amylase, human α -hemoglobin or human β -hemoglobin, is inserted adjacent the promoter of the TMV coat protein coding sequence, which has been deleted. In another example, a tyrosinase coding sequence such as isolated from I Streptomyces antibioticus is inserted adjacent the same promoter of TMV, oat mosaic virus (OMV) or rice necrosis virus (RVN). Recombinant virus can be prepared as described above, using the resulting recombinant plant viral nucleic acid. Tobacco or germinating barley is infected with the recombinant virus or recombinant plant viral nucleic acid. The viral nucleic acid self-replicates in the plant tissue to produce the enzymes amylase or tyrosinase. The activity of this tyrosinase leads to the

production of melanin. See, for example, Huber, M. et al. (36).

In a further example, a cyclodextrin glucanotransferase coding sequence, such as isolated from Bacillus sp. No. 17-1 (see U.S. Patent 4,135,977) is inserted adjacent the promoter of the viral coat protein of a nucleotide sequence derived from OMV, RNV, PVY or PVX in which the coat protein coding sequence has been removed, and which then contains a non-native promoter and coat protein gene. Corn or potato is infected with the appropriate recombinant virus or recombinant plant viral nucleic acid to produce the enzyme cyclodextrin glucotransferase. The activity of this enzyme leads to the production of cyclodextrin, which is useful as a flavorant or for drug delivery.

In some plants, the production of anti-sense RNA as a product can be useful to prevent the expression of certain phenotypic traits. Particularly, some plants produce substances which are abused as drugs (e. g., cocaine is derived from the coca plant, and tetrahydrocannabinol (THC) is the active substance of abuse derived from cannabis or marijuana plants). An anti-sense RNA complementary to the plant RNA necessary for the production of an abusable substance would prevent the production of the substance. This could prove to be an effective tool in reducing the supply of illegal drugs.

A still further feature of the invention is a process for the production of an enzyme suitable for the stereospecific catalysis of an organic compound. The non-native nucleic acid comprises the transcribable sequence which leads to the production of the desired product. This process involves the infection of the appropriate host with a recombinant

virus or recombinant plant viral nucleic acid such as those described above, the growth of the infected host to produce the desired product and the isolation of the desired product. The growth of the infected host is in accordance with conventional techniques, as is the isolation of the resultant product. The stereospecific enzyme is then utilized to catalyze the desired reaction. One use of stereospecific enzymes is in the separation of racemate mixtures.

In one example, a suitable esterase or lipase coding sequence such as isolated from an appropriate microorganism is inserted adjacent the promoter of the viral coat protein of a nucleotide sequence derived from TMV, oat mosaic virus (OMV) or rice necrosis virus (RNV) in which the coat protein coding sequence has been removed and which then contains a non-native promoter and coat protein gene. Tobacco or germinating barley is infected with the recombinant virus or recombinant plant viral nucleic acid to produce the esterase or lipase enzyme. This enzyme is isolated and used in the stereospecific preparation of a compound such as naproxen, as described in EP-A 0233656 or EP-A 0227078.

I An esterase coding sequence is isolated from the appropriate microorganism, such as Bacillus subtilis, Bacillus licheniformis (a sample of this species is deposited with the American Type Culture Collection, Rockville, Maryland (ATCC) under Accession No.

I 11945), Pseudomonas fluorescens, Pseudomonas putida (a sample of this species is deposited with the Institute for Fermentation (IFO), Osaka, Japan, under

I Accession No. 12996), Pseudomonas riboflavina (a sample of this species is deposited with IFO under

I Accession No. 13584), Pseudomonas ovalis (a sample of this species is deposited with the Institute of

I Applied Microbiology (SAM), University of Tokyo, Japan, under Accession No. 1049), Pseudomonas aeruainosa (IFO 13130), Mucor angulimacrosporus (SAM 6149), Arthrobacter paraffineus (ATCC 21218), Strain is III-25 (CBS 666.86), Strain LK 3-4 (CBS 667.86), Strain Sp 4 (CBS 668.86), Strain Thai III 18-1 (CBS 669.86), and Strain Thai VI 12 (CBS 670.86).

I Advantageously, cultures of species Bacillus subtilis include cultures of species Bacillus species Thai 1-8 (CBS 679.85), species Bacillus species In IV-8 (CBS 680.85), species Bacillus species Nap 10-M (CBS 805.85), species Bacillus species Sp 111-4 (CBS 806.85), Bacillus subtilis 1-85 (Yuki, S. et al., Japan J. Gen. 42:251 (1967)), Bacillus subtilis 1-85/pNAPT-7 (CBS 673.86), Bacillus subtilis 1A-40/pNAPT-8 (CBS 674.86), and Bacillus subtilis 1A-40/pNAPT-7 (CBS 675.86). Advantageously, cultures of Pseudomonas fluorescens include a culture of species Pseudomonas species Kpr 1-6 (CBS 807.85), and Pseudomonas fluorescens species (IFO 3081).

A lipase coding sequence is isolated from the appropriate microorganism such as the genera Candida, Rhizopus, Mucor, Aspergilus, Penicillium, Pseudomonas, Chromobacterium, and Geotrichum. Particularly preferred is the lipase of Candida cylindracea (Qu-Ming et al., Tetrahedron Letts. 27, 7 (1986)).

A fusion protein can be formed by incorporation of the non-native nucleic acid into a structural gene of the viral nucleic acid, e.g., the coat protein gene. The regulation sites on the viral structural gene remain functional. Thus, protein synthesis can occur in the usual way, from the starting codon for methionine to the stop codon on the foreign gene, to produce the fusion protein. The fusion protein

contains at the amino terminal end a part or all of the viral structural protein, and contains at the carboxy terminal end the desired material, e.g., a stereospecific enzyme. For its subsequent use, the stereospecific enzyme must first be processed by a specific cleavage from this fusion protein and then further purified. A reaction with cyanogen bromide leads to a cleavage of the peptide sequence at the carboxy end of methionine residues (5.0. Needleman, "Protein Sequence Determination", Springer Publishers, 1970, N.Y.). Accordingly, it is necessary for this purpose that the second sequence contain an additional codon for methionine, whereby a methionine residue is disposed between the N-terminal native protein sequence and the C-terminal foreign protein of the fusion protein. However, this method fails if other methionine residues are present in the desired protein. Additionally, the cleavage with cyanogen bromide has the disadvantage of evoking secondary reactions at various other amino acids.

Alternatively, an oligonucleotide segment, referred to as a "linker," may be placed between the second sequence and the viral sequence. The linker codes for an amino acid sequence of the extended specific cleavage site of a proteolytic enzyme as well as a specific cleavage site (see, for example, U.S. Patent Nos. 4,769,326 and 4,543,329). The use of linkers in the fusion protein at the amino terminal end of the non-native protein avoids the secondary reactions inherent in cyanogen bromide cleavage by a selective enzymatic hydrolysis. An example of such a linker is a tetrapeptide of the general formula Pro-Xaa-Gly-Pro(SEQ ID NO: 1) (amino-terminal end of non-native protein), wherein Xaa is any desired amino acid. The overall cleavage is

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effected by first selectively cleaving the xaa-Gly bond with a collagenase (E.C. 3.4.24.3., Clostridiopeptidase A) then removing the glycine residue with an aminoacyl-proline aminopeptidase (aminopeptidase-P, E.C. 3.4.11.9.) and removing the proline residue with a proline amino peptidase (E.C. 3.4.11.5). In the alternative, the aminopeptidase enzyme can be replaced by postproline dipeptidylaminopeptidase. Other linkers and appropriate enzymes are set forth in U.S. Patent No. 4,769,326.

A still further feature of the invention is a process for the induction of male sterility in plant. Male sterility can be induced by several mechanisms, including, but not limited to, an anti-sense RNA mechanism, a ribozyme mechanism, or a protein mechanism which may induce male sterility or self-incompatibility or interfere with normal gametophytic development. The second nucleotide sequence of the chimeric nucleotide sequence comprises the transcribable sequence which leads to the induction of male sterility. This process involves the infection of the appropriate plant with a virus, such as those described above, and the growth of the infected plant to produce the desired male sterility. The growth of the infected plant is in accordance with conventional techniques.

Male sterility can be induced in plants by many mechanisms including, but not limited to (a) absence of pollen formation, (b) formation of infertile and/or non-functional pollen, (c) self-incompatibility, (d) inhibition of self-compatibility, (e) perturbation of mitochondrial function(s), (f) alteration of the production of a hormone or other biomolecule to interfere with normal

gametophytic development, or (g) inhibition of a developmental gene necessary for normal male gametophytic tissue. These mechanisms may be accomplished by using anti-sense RNA, ribozymes, genes or protein products. The recombinant plant viral nucleic acids of the present invention contain one or more nucleotide sequences which function to induce male sterility in plants. To accomplish this function, the recombinant plant viral nucleic acids may contain a nucleotide sequence, a single gene or a series of genes.

Male sterility traits could be formed by isolating a nuclear-encoded male sterility gene. Many of these genes are known to be single genes. For example, Tanksley et al. (37) placed ms-10 in CIS with a rare allele of the tightly linked enzyme-coding gene Prx-2. The Prx-2 allele is codominant, allowing selection for heterozygous plants carrying the recessive ms-10 allele in backcross populations and eliminating the need for progeny testing during transfer of the gene into parents for hybrid production. A male-sterile anthocyaninless plant (ms-10 aa/ms-10aa) was crossed to a heterozygous, fertile plant in which a rare peroxidase allele was in cis with the recessive male-sterile allele (ms-10 Prx-2'/+Prx-2+). Male sterile plants were selected from the progeny (ms-10 Prx-2'/ms-10aa). Once the male-sterile gene has been transferred into a prospective parental line, sterile plants can be selected at the seedling stage either from backcross or F₂ seed lots.

In pearl millet, recessive male sterile genes were found in vg 272 and IP 482. Male sterility in pearl millet line Vg 272 and in IP 482 is essentially controlled by a single recessive gene. Male

sterility in Vg 272 is due to a recessive gene, ms, which has no effect on meiosis in pollen mother cells, but acts after separation of microspores from tetrads but before onset of the first mitotic division.

Dewey et al. (39) isolated and characterized a 3547 bp fragment from male sterile (cms-T) maize mitochondria, designated TURF 243. TURF 243 contains two long open reading frames that could encode polypeptides of 12,961 Mr and 24,675 Mr. TURF 243 transcripts appeared to be uniquely altered in cms-T plants restored to fertility by the nuclear restorer genes Rf1 and Rf2. A fragment of maize mtDNA from T cytoplasm was characterized by nucleotide sequence analysis. To obtain isolation of nucleic acids, mitochondrial RNA (mtRNA), and mtDNA were prepared from six- to seven-day-old dark grown seedlings of Zea Mays L. by conventional techniques.

Another means by which male sterile traits could be formed is by the isolation of a male sterility gene from a virus. There are several viruses or virus-like particles that induce male sterility in plants. Recent work suggests that viroid-like agents in male sterile beets may occur. (40). Cytoplasmic male sterility may be conditioned by a discrete particle such as a plasmid or an inclusion. Viruses are not seed transmitted with the regularity of cytoplasmonic systems. Viroids can be transmitted through pollen. Transfer of a factor of some kind across a graft union has been demonstrated in petunia, beet, sunflower, and alfalfa. There is no direct effect on the fertility of the scion, but selfs or crosses by a maintainer on the grafted scion produced male sterile plants in the next generation. Cms beets grown at 36°C for 6 weeks, then at 25°C,

produced fertile plants from new shoots possibly due to elimination of "cytoplasmic spherical bodies", but progenies from the plants reverted to sterility after three generations at normal growing conditions.

Cytoplasmic male sterility in the broad bean plant I (*Vicia faba*) was found to be caused by the presence of virus or virus-like particles. Possibly a case similar to a cms-system occurs in garlic. Pollen degeneration typical of sporophytic cms plants was found, but electron microscope studies showed rickettsia-like inclusions in the anthers, which could be eliminated with antibiotics, causing the pollen to become fertile (41).

Male sterile traits could be formed by a third method of introducing an altered protein, using a transit peptide sequence so that it will be transported into the mitochondria, and perturbing the mitochondrial functions. This protein could work to overwhelm normal mitochondrial function or reduce a metabolite required in a vital pathway. It is widely believed that slight perturbations in the mitochondria will lead to male sterility. Remy et al. (42) conducted a two dimensional analysis of chloroplast proteins from normal and cytoplasmic

I male-sterile B. napus lines. Chloroplast and mitochondrial DNAs of N and cms lines of B. napus were characterized and compared using restriction enzyme analysis. Identical restriction patterns were

I found for chloroplastic DNAs from the cms B. napus lines and the cms lines of the Japanese radish used

I₃ to transfer the cms trait into B. napus. In Remy's study, chloroplast proteins from stroma and

I thylakoids of N and cms lines of B. napus were characterized and compared using a 2-D polyacrylamide gel separation. It was shown that (1) stromal

(el) compartments of the two lines were very similar, and (2) the lines could be distinguished by the spots corresponding to the β subunits of coupling factor CP, from the ATPase complex.

(13) A fourth method for inducing male sterility in plants is by inducing or inhibiting a hormone that will alter normal gametophytic development -- for example, inhibiting the production of gibberellic acid prior to or at the flowering stage to disturb pollen formation, or modifying production of ethylene prior to or at the flowering stage to alter flower formation and/or sex expression.

A fifth method for inducing male sterility in plants is by inhibiting a developmental gene required for the normal male gametophytic tissue, for example, using anti-sense RNA that is complementary to the developmental signal RNA or mRNA. Padmaja et al. (43) discusses cytogenetical investigations on a spontaneous male-sterile mutant isolated from the Petunia inbred lines. Male sterility was found to be associated with atypical behavior of tapetum, characterized by prolonged nuclear divisions and untimely degeneration as a result of conversion from glandular to periplasmoidal type.

A sixth method for inducing male sterility in plants is by isolating a self-incompatibility gene and using the gene in the vector of the present invention. Self-incompatibility (S) gene systems that encourage out-breeding are present in more than 50% of the angiosperm plant families (44). Multiple S gene systems are known in some species. In several systems, abundant style glycoproteins (S glycoproteins) have been identified. These glycoproteins are polymorphic and can be correlated with identified S alleles. S genes, corresponding to

I the style glycoproteins of N. alaba and B. oleraceae have been cloned and sequenced. Amino acid substitutions and deletions/insertions, although present throughout the sequences, tend to be clustered in regions of hypervariability that are likely to encode allelic specificity.

A seventh method for inducing male sterility in plants is by blocking self incompatibility, by the engineering of a protein that will bind and inactivate the compatibility site or by turning off self-compatibility, by the engineering of an anti-sense RNA that will bind with the mRNA to a self-compatibility protein.

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Specific effects resulting in male sterility can range from the early stages of sporogenous cell formation right through to a condition in which anthers containing viable pollen do not dehisce. Some or all of the developmental stages within this range may be affected. Some of the more obvious specific effects include, the following examples:

1) Meiosis is disrupted, leading to degeneration of the pollen mother cells or early microspores in which case pollen aborts and anther development is arrested at an early stage.

2) Exine formation is disrupted and microspores are thin-walled, perhaps distorted in shape, and nonviable. Anthers are generally more developed than the exines, but still not normal.

3) Microspore vacuole abnormalities, decreased starch deposition and tapetum persistence are evident. Pollen is nonviable and anthers are still not normal.

4) Pollen is present and viable, and anthers appear normal but either do not dehisce or show much delayed dehiscence.

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5) Self incompatibility mechanisms disrupt or prevent enzymatic digestion of the style by the pollen grain.

Male sterility in plants may be induced by the mechanisms listed above at any stage prior to pollen shed. The male sterility mechanism selected may be applied to plants in the field (or in the greenhouse) at any time after seedling emergence and before pollen shed. The exact time of application will depend on the male sterility mechanism used and the optimum effectiveness in producing male sterile plants.

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EXAMPLES

In the following examples, enzyme reactions were conducted in accordance with manufacturers recommended procedures, unless otherwise indicated. Standard techniques, such as those described in Molecular Cloning (7), Meth. in Enzymol. (9) and DNA Cloning (8), were utilized for vector constructions and transformation unless otherwise specified.

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COMPARATIVE EXAMPLES

The following comparative examples demonstrate either the instability of prior art recombinant viral nucleic acid during systemic infection of host plants or the inability to systemically infect plants and to efficiently produce the product of the inserted nonnative gene.

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Comparative Example 1

Recombinant plant viral nucleic acid was prepared by inserting the chloramphenical acetyltransferase (CAT) gene which had been fused behind a TMV subgenomic RNA promoter between the 30K

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and coat protein genes of TMV. pTMV-CAT-CP was prepared as described by Dawson, W.O. et al. (11). Briefly, pTMV-CAT-CP was constructed by cutting pTMV204, a full-genomic cDNA clone of TMV strain U1 (4) with NcoI (nt. 5460), blunting with Klenow fragment of DNA polymerase I, adding PstI linkers (CCTGCACG from Boehringer-Mannheim Biochemicals), excising with PstI and NsiI (nt. 6207), and ligating this 747-bp fragment into the NsiI site (nt. 6207) of pTMV-S3-CAT-28, a modified TMV with the CAT ORF substituted for the coat protein ORF (45). TMV nucleotide numbering is that of Goelet et al. (46). Correct ligation and orientation of each construct were checked by restriction mapping and sequencing.

Inoculations. In vitro transcription of plasmid DNA constructs and inoculation procedures were as described previously (3). Virus was propagated systemically in Xanthi tobacco (Nicotiana tabacum L.) and Nicotiana sylvestris: Xanthi-nc tobacco was used as a local lesion host. Plants were grown in a greenhouse prior to inoculations and then subsequently maintained in plant growth chambers at 25° with a 16-hour photoperiod of approximately 2000 lx.

CAT Assays. Amounts of CAT activity were assayed essentially by the procedures described (47), 200 mg of leaf tissue were macerated in assay buffer followed by addition of 0.5 mM acetyl CoA and 0.1 μ Ci [14 C]-chloramphenicol, incubation for 45 minutes at 37°, extraction and resolution by thin-layer chromatography, and finally autoradiography.

RNA Analysis. Four days after inoculation, total RNA from infected leaves was extracted as described (47a). For blot hybridization analysis, RNA was electrophoresed in 1.2% agarose gels,

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transferred to nitrocellulose, and hybridized with
nick-translated cDNA of TMV (nts. 5080-6395) in
pUC119 or pCM1 (Pharmacia) which contains the CAT
ORF. Total RNA from infected leaves also was
analyzed by RNase protection assays for wild-type
sequences essentially as described in Ausubel et al.
40 (48). The 3' half (BamHI:nt. 3332-PstI:nt. 6401) of
pTMV204 was cloned into pT7/T3-19 (from BRL). After
H EcoRI digestion (nt. 4254), ³²P-labeled transcripts
complementary to the 3' viral sequences were produced
with T7 RNA polymerase. An excess amount of the
probe was hybridized to RNA samples, treated with 40
82 μ g/ml RNase A (Sigma) and 300 U RNase T1 (BRL)
extracted, denatured with DMSO and glyoxal, and
electrophoresed in 1.2% agarose gels which were
subsequently dried and exposed to Kodak X-ray film.

Construction of cDNA Clones of ProgenY Virus.

RNA was extracted from purified virions and cDNA was
prepared as previously described (4). Double-stranded
cDNA was digested with BamHI (nt. 3332) and SacI (nt.
6142) and cloned into BamHI- and SacI-digested pUC19.
Nucleotide sequencing of DNA was by the
dideoxynucleotide chain terminating procedure (49).

Results. In vitro transcripts of pTMC-CAT-CP,
which had the CAT cartridge inserted upstream of the
coat protein gene, resulted in CAT-CP, a hybrid virus
7452 nucleotides in length and a gene order of 126K,
183K, 30K, CAT and coat protein. In vitro
I transcripts were used to inoculate leaves of N.
tabacum L. varieties Xanthi and Xanthi-nc and N.
sylvestris. Results were compared to those from
plants infected with wild-type virus, TMV 204, or the
free-RNA virus, S30CAT-28, that expresses CAT as a
replacement for coat protein (45). CAT-CP replicated
effectively and moved from cell to cell in inoculated

I leaves similarly to TMV 204. Necrotic lesions developed on Xanthi-nc tobacco at approximately the same time and were of the same size as those caused by TMV 204 and S3-CAT-2B. CAT-CP induced no symptoms in inoculated leaves of the systemic hosts, Xanthi tobacco and N. sylvestris, but produced mosaic symptoms in developing leaves similar to those produced by TMV 204. The concentration of virions in cells infected with CAT-CP, estimated by yields obtained after virion purification and by transmission electron microscopy of thin sections of inoculated leaves, appeared to be approximately equal to that from a TMV 204 infection.

CAT-CP is 7452 nucleotides long, compared to 6395 nucleotides for TMV 204, which would result in CAT-CP virions 350 nm in length, compared to the 300 nm virions of wild-type TMV. Virus was purified from inoculated leaves of CAT-CP-infected plants and analyzed by transmission electron microscopy. Most of the virions from the CAT-CP infections were 350 nm in length. One problem in assessing the length of virions of TMV UI viewed by electron microscopy is that preparations normally contain fragmented and end-to-end aggregated virions in addition to individual genomic-length virions. To determine the proportion of 350- to 300-nm virions, distinct, individual virions of each size were counted. The ratio of 350/300 nm virions in leaves inoculated with CAT-CP was 191:21, compared to 12:253 from the wild-type infection. The 350-nm virions in wild-type TMV infection probably resulted from the end-to-end aggregation of fragmented virions, since TMV UI has a propensity to aggregate end-to-end and all length virions can be found. These data suggest that the

extra gene of CAT-CP was maintained and encapsidated in these inoculated leaves.

CAT activity was detected in leaves inoculated with CAT-CP using in vitro RNA transcripts or the subsequent first or second passage local lesions. From more than one hundred samples assayed, a range of variation was found among different positive samples. Similar levels of CAT were found in CAT-CP-infected leaves as those infected with the coat protein-less mutant, S3-CAT-2 B. Only background amounts were detected in TMV 204-infected or healthy leaves.

The host range of CAT-CP was compared to that of wild-type TMV by inoculating a series of hosts known to support replication of TMV and by screening for CAT activity. CAT activity was detected in inoculated leaves of Zinnia elegans Jacq., Lunaria annua L., Beta vulgaris L., Calendula officinalis L., and Spinacia oleracea L., which represent three plant families in addition to the Solanaceae. This indicated that this alteration of the TMV genome did not appear to alter the host range.

In order to determine whether CAT-CP produced an additional subgenomic RNA as a result of the inserted sequences, total RNA from infected leaves was extracted and compared to that of wild-type TMV by blot hybridization analysis, using a TMV or a CAT DNA probe. Xanthi tobacco leaves infected with CAT-CP previously passaged twice in xanthi-nc tobacco were chosen because they contained a population of CAT-CP and progeny virus with deletions to be compared to wild-type TMV. Two distinct genomic RNAs were detected. The largest hybridized to both TMV and CAT probes, whereas the smaller genomic RNA hybridized only to the TMV probe and comigrated with wild-type

Tv genomic RNA. Three distinct, small RNAs were found in RNA from CAT-CP-infected leaves, compared to two from TMV 204-infected leaves. The smaller RNAs that comigrated with the subgenomic messages for the coat and 30K proteins of wild-type TMV hybridized only to the Tv-specific probe. A larger subgenomic RNA from CAT-CP-infected leaves hybridized to both the CAT and TMV probes. Assuming that as for the subgenomic mRNAs of wild-type TMV, this larger subgenomic RNA is 3' coterminous with the genomic RNA (50), these results are consistent with the extra CAT-CP mRNA predicted for expression of CAT. The putative CAT-CP subgenomic RNA for 30K protein, containing the 30K, CAT, and coat protein ORFs was not observed, possibly because bands in the region between 2.4 and 4.4 kb were obscured by viral RNAs adhering during electrophoresis to host rRNAs and were difficult to resolve (50, 51).

The amounts of CAT activity in upper, systemically infected leaves were variable and much lower than in inoculated leaves, and in many cases none was detected. Hybridizations with Tv and CAT probes demonstrated that the proportion of virus-retaining CAT sequences was quickly reduced to undetectable levels. The transition from CAT-CP to a population of virus with the inserted CAT ORF deleted occurred during systemic invasion of the plant and sometimes in inoculated leaves. In contrast, CAT sequences and CAT activity often were detected in leaves inoculated with virus that had been passaged through single lesions three or four times.

CAT-CP virions were examined from systemically infected Xanthi tobacco leaves approximately 30 days after inoculation. Quantification of virions from

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the uppermost leaves of the plants infected with CAT- CP produced a ratio of 350- /300-nm virions of 78:716. This was compared to a ratio of 191:21 in inoculated leaves, indicating that the major component of the population shifted to 300-nm virions during systemic infection. The deleted progeny virus recovered after continued replication of CAT-CP was identical in host range and symptomatology to wild-type TMV.

cDNA of the region that encompassed the CAT insertion (nts. 3332-6142) was cloned from the progeny CAT-CP virion RNA from systemically infected Xanthi leaves to sample the virus population. Characterization of nine cDNA clones by size and restriction mapping indicated that eight were identical with wild-type TMV.

One cDNA clone appeared to be the size predicted for the CAT-CP construct, but the restriction map varied from that predicted for CAT-CP. Five clones that were evaluated by size and restriction analysis as wild-type were sequenced through the region of the CAT insertion and also through a portion of the coat protein gene, and found to be identical to the parental wild-type virus. This suggested the inserted sequences could be excised, giving rise to wild-type TMV.

To corroborate this possible excision, samples of the total leaf RNA used in the blot hybridization analysis were analyzed by RNase protection assays using T7-produced minus-strand RNA complementary to in inoculated leaves. In contrast, CAT sequences and CAT activity often were detected in leaves inoculated with virus that had been passaged through single lesions three or four times.

CAT-CP virions were examined from systemically infected Xanthi tobacco leaves approximately 30 days after inoculation. Quantification of virions from the uppermost leaves of the plants infected with CAT-CP produced a ratio of 350- /300-nm virions of 78:716. This was compared to a ratio of 191:21 in inoculated leaves, indicating that the major component of the population shifted to 300-nm virions during systemic infection. The deleted progeny virus recovered after continued replication of CAT-CP was identical in host range and symptomatology to wild-type TMV.

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To corroborate this possible excision, samples of the total leaf RNA used in the blot hybridization analysis were analyzed by RNase protection assays using T7-produced minus-strand RNA complementary to nucleotides 4254-6395 of wild-type TMV. The presence of wild-type sequences in this region would result in

a protected RNA of 2140 nucleotides. A band this size from the CAT-CP RNAs comigrated with a similar band produced using wild-type RNA to protect the probe. These data confirmed that the inserted sequences of CAT-CP could be precisely deleted. Taking into consideration the presence of repeated sequences in CAT-CP RNA that allow the bulge loop in the hybrid between CAT-CP and the wild-type TMV probe RNA to occur over a range of positions within the repeats, the RNase protection of wild-type probe by CAT-CP RNA should produce sets of bands that would fall within two nucleotide size ranges, 683-935 and 1202-1458. The other two major bands seen are of these sizes, corroborating the presence of CAT-CP RNA in these samples.

The loss of the inserted sequences of CAT-CP appeared to be due to two sequential processes. First was the loss of inserted sequences in individual molecules, as shown by the sequence analysis of cDNA clones of progeny virus. Since the deletion occurred between repeated sequences, it is possible that this occurred by homologous recombination as described for other plus-sense RNA viruses (52-54). The second process resulted in a selected shift in the virus population. The RNase protection assays, in which the virus population was sampled, demonstrated that both CAT-CP and wild-type virus could be components of the population in inoculated leaves. The lack of CAT-CP in systemically infected leaves was probably due to a shift in the virus population, possibly because the original hybrid could not effectively compete with the deleted progeny wild-type virus in terms of replication and systemic movement.

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Comparative Example 2

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A recombinant plant viral nucleic acid was prepared by inserting the CAT gene which had been fused behind a TMV subgenomic RNA promoter between the coat protein gene and the nontranslated 3' region of TMV. pTMV-CP-CAT was prepared as described by Dawson et al. (II) Briefly, pTMV-CP-CAT was constructed by cutting pTMV-S3-CAT-28 with HindIII (nt. 5081), blunting with Klenow fragment of DNA polymerase I, adding PstI and NsiI (nt. 6207), and ligating this 1434-bp fragment in the NsiI site (nt. 6207) of pTMV204. Correct ligation and orientation of each construct were checked by restriction mapping and sequencing.

Plant inoculations, CAT assays, RNA analysis and construction of cDNA clones of progeny were performed as described in Comparative Example I. pTMV-CP-CAT, the larger hybrid virus construct, contained a 628-nucleotide repeat of that portion of the 30K gene containing the coat protein subgenomic promoter and the origin of assembly. This construct should produce a virus, CP-CAT, 7822 nt long with a gene order of 126K, 183K, 30K, coat protein, and CAT. CP-CAT replicated poorly. It produced necrotic lesions in Xanthi-nc that were small, approximately one-half the diameter of wild-type virus lesions, and their appearance was delayed by two days. Transmissibility of CP-CAT from these lesions was at a level approximately one-hundredth that of CAT-CP or wild-type TMV. No systemic symptoms appeared in Xanthi or N. sylvestris plants and the virus infection was transferrable only from inoculated leaves. Low but reproducible levels of CAT activity were found in CP-CAT-infected leaves. Since the replication of this

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chimeric virus was so impaired, characterization did not proceed any further.

In contrast to CAT-CP, when CP-CAT was allowed to replicate for extended periods in the systemic hosts, no wild-type-like virus symptoms ever were observed in upper leaves of plants and virus was never recovered from them, suggesting that this hybrid virus did not delete the inserted sequences in a manner to create a wild-type-like virus.

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Comparative Example 3

A full-length DNA copy of the TMV genome is prepared and inserted into the PSTI site of pBR322 as described by Dawson, W.O. et al. (t). The viral coat protein gene is located at position 5711 of the TMV genome adjacent the 30k protein gene. The vector containing the DNA copy of the TMV genome is digested with the appropriate restriction enzymes and exonucleases to delete the coat protein coding sequence. For example, the coat protein coding sequence removed by partial digestion with ClaI and NsiI, followed by religation to reattach the 3'-tail of the virus. Alternatively, the vector is cut at the 3' end of the viral nucleic acid. The viral DNA is removed by digestion with Bal31 or exonuclease III up through the start codon of the coat protein coding sequence. A synthetic DNA sequence containing the sequence of the viral 3'-tail is then ligated to the remaining 5'-end. The deletion of the coding sequence for the viral coat protein is confirmed by isolating TMV RNA and using it to infect tobacco plants. The isolated TMV RNA is found to be non-infective under natural conditions.

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The 314-bp Sau3A fragment (NH₂ terminus of the Tn5 NPTII gene) from pNEO was filled in with Klenow

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polymerase and ligated to SalI ($pd[GGTCGACC]$) linkers. It was then digested with SalI and PstI and inserted into PstI/SalI-digested pUC128 (55) to give pNU10. The pNEO plasmid was digested with AsuII, filled in with Klenow polymerase and ligated to XhoI linkers ($pd[CCTCCGAGG]$) to give pNX1. The pNX1 was digested with XhoI, filled in with Klenow polymerase, digested with PstI and ligated into PstI/SmaI-digested pNU10 to give pNU116.

The XhoI/SalI fragment from pNU116 (NPTII sequences) is ligated adjacent the coat protein promoter. The resultant RFVNA containing the NPTII gene insert was applied to twelve Nicotiana tabacum (cv. Xanthi-NC), a cultivar that has been backcrossed to contain the N gene for TMV resistance and to twelve N. tabacum (cv. Xanthi), a cultivar that does not contain the N gene. In both tobacco cultivars, no systemic spread was observed in any inoculated plant. The N. tabacum (cv. Xanthi NC) showed the characteristic flecking spots on the inoculate leaf indicating resistance to the virus. The N. tabacum (cv. Xanthi) exhibited no flecking or systemic symptoms.

CL Comparative Example 4

P A recombinant plant viral nucleic acid containing the NFTII coding sequence was prepared as described in Comparative Examples 1 and 3. The NFTII and coat protein coding sequences were each adjacent an "O" coat protein promoter. The presence of the coat protein gene should render the vector capable of being systemically spread.

I The resultant RFVNA containing the NPTII-
L inserted gene was inoculated on twelve N. tabacum (cv. Xanthi NC) and twelve N. tabacum (cv. Xanthi NC)

I showed the flecking in each of the twelve plants, as with Comparative Example 1. The N. tabacum (cv. Xanthi) plants showed systemic spread of the vector in all twelve plants.

I Leaf discs from N. tabacum (cv. Xanthi) leaves were cultured on media containing kanamycin. None of the tissue survived in culture, indicating a loss or dysfunction of the NFTII gene. Subsequent electron photomicroscopy of the present vector containing the NFTII gene recovered from the leaves of treated N. tabacum (cv. Xanthi) plants showed that the present vector had lost a section of the vector corresponding to the NPTII gene, indicating a breakage and recombination of the vector.

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EXAMPLES OF THE PREFERRED EMBODIMENTS

The following examples further illustrate the present invention. These examples are intended merely to be illustrative of the present invention and are not to be construed as being limited.

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EXAMPLE 1

I Construction of Bacterial Plasmids. Numbers in parentheses refer to the TMV-U1 sequence (46). DNA manipulations were performed essentially as described in (48). All plasmids were propagated in E. coli strain JM109 except for pTBN62 (DH5 α ; Gibco BRL; and H8101).

pTKU1 (Fia. 1). The 7.3 kb pTMV204 (4) PstI fragment (TMV-U1 genome and λ phage promoter from pPM1 (3) was subcloned into pUC19 to give pTP5. pTMV204 ApaI fragment (5455-6389) was ligated to oligonucleotides pd[CAGGTACCC] and d[GGGTACCTGGGCC], (SEQ ID No: 2), digested with KpnI (underlined within nucleotide sequence) and NcoI (5459) and ligated into

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NcoI/KpnI digested pTP5 to produce pTPK10. pTKU1 was constructed by subcloning the 7.3 kb PstI/KpnI fragment from pTPK10 into PstI/KpnI-digested pUC118. pTKU1 contained a DNA copy of the entire TMV-VI genome downstream of the λ phage promoter from pPML. KpnI digestion and in vitro transcription of pTKU1 gave infectious TMV RNA. pTKU1 was constructed because PstI sites in the odontoglossum ring spot virus (ORSV, sometimes referred to as TMV-O) coat protein, DHFR and NFTII ORFs prohibited the use of this restriction enzyme (employed to linearize pTMV204; 4) to digest plasmid DNA of the hybrid constructs and produce infectious in vitro transcripts.

pT82 (Fig. 1). pTMVS3-28 (45) was a derivative of pTMV204 in which the coat protein initiation codon was mutated to ACG and a XhoI site replaced the entire coat protein coding sequence. The 1.9 kb NcoI/SalI fragment (5459-SalI site in p8R322) from pTMVS3-28 was ligated into NcoI/SalI-digested pNEO (56) to give pNS283. pBabsI was a 2.4 kb EcoRI cDNA clone from ORSV virion RNA with nucleotide, ORF and amino acid sequence similarities to TMV-UI (nts 4254-6370). A 680 bp pBabsI HincII/EarI (Klenow polymerase infilled) fragment (containing the ORSV coat protein ORF and 203 bases upstream of its AUG) was ligated into the NstI site (6202; blunt-ended with T4 DNA polymerase) of pNS283 to produce pB31. The NcoI/SalI fragment from p831 was then ligated into the NcoI/SalI-digested pTMV204 (replacing the corresponding wild-type fragment 5459-SalI site in pBR322) to give pTB281. pTB2 was constructed by ligating the BamHI/SplI fragment from pTB281 into BamHI/SplI-digested pTKU1 (replacing the corresponding wild-type fragment 3332-6245).

pNC4X (57). pNC4X consisted of the R67 DHFR gene cloned into pUC8X. The plasmid contained a XhoI site eight bases upstream of the initiation codon for the DHFR gene. In addition, the stop codon and five bases of carboxy-terminal DHFR sequence were deleted and replaced by a SalI site.

H pNU116. A 315 bp pNEO Sau3S (Klenow polymerase infilled) fragment (NH_2 terminus of Tn5 NPTII gene) was ligated to SalI ($\text{pd}[\text{GGTCGACC}]$) linkers, SalI/FstI digested, and inserted into FstI/SalI-digested pUC128 (55) to give pNU10. pNEO was digested with AsuII, infilled with Klenow polymerase and ligated to XhoI linkers ($\text{pd}[\text{CCTCGAGG}]$) to generate pNX1. pNU116 was constructed by digesting pNX1 with XhoI, infilling with Klenow polymerase, digesting with PstI and ligating the resulting 632 bp fragment (COOH terminus of the Tn5 NPTII gene) into PstI/SmaI-digested pNU10. This manipulation of the NPTII gene removed an additional ATG codon 16 bases upstream of the initiation codon, the presence of which decreased NPTII activity in transformed plant cells (58).

pTBD4 and pTBN62 (Fia. 1). XhoI/SalI fragments from pNC4X (DHFR sequence) and pNU116 (NPTII sequence) respectively were ligated into the XhoI site of pT82 in the same sense as the TMV coding sequences.

In Vitro Transcription and Inoculation of Plants. Plants grown as in (45) were inoculated with in vitro transcripts TB2 (nt. 6602), T8D4 (nt. 6840) and TBN62 (nt. 7434) from KpnI digested pTBD2, pTBD4 and pTBN62, respectively. The in vitro transcription method was as previously described.

Analysis of Progeny Virion RNA. Virus purification was essentially as described by Gooding and Hebert (59) with one precipitation with

14,33 polyethylene glycol (8% PEG, 0.1M NaCl; 0°C 1 hr) and one ultracentrifugation (151,000-235,000 x g; 90 min). Virion RNA was extracted by digesting 1 mg virus with 0.2 µg Froteinase K in 10mM Tris HCl, pH 7.5, 1mM EDTA, 0.1% SDS at 37°C for 1 hr, followed by phenol/chloroform extractions. RNA samples were DMSO-denatured, glyoxalated, electrophoresed in 1% agarose gels and transferred to nitrocellulose (pore size 0.45 µm; Schleicher and Schull; 48). The transfers were probed with [α -³⁵S]-dATP (New England Nuclear) labelled (50) restriction fragments. RNase protection assays were as described in (48). TBD4-38 and pTBN62-38 contained BamHI/KpnI fragments (nts. 3332-6396) from pTBD4 and pTBN62, respectively, cloned into BamHI/KpnI-digested pBluescript SKI- (Stratagene)

P Immunological Detection of NPTII. Sample preparation and Western analysis were as described previously (45). Leaf samples were ground in liquid N₂ and extraction buffer (10% glycerol, 62.5mM Tris HCl pH 7, 5% mercaptoethanol, 5mM phenylmethylsulfonyl fluoride). Equivalent protein concentrations were determined and absolute concentrations estimated by Bradford assay (Stratagene; 61), with bovine serum albumin as standard. Western transfers were probed with antiserum to NPTII (1:500; 5 Prime, 3 Prime, Inc.) and then with alkaline phosphatase-conjugated goat anti-rabbit IgG (1:1000).

NFIII Activity Assays. NPTII activity was detected by its phosphorylation of neomycin sulphate.

Enzyme assays were as described in (62) except the extraction buffer was as described above and dilution series of purified NPTII (5 Prime, 3 Prime, Inc.) in healthy tissue were included.

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Leaf Disc Assays to Screen for Resistance to Kanamycin Sulphate. NPTII confers resistance to the aminoglycoside kanamycin (56). Young systemic leaves 12 days post-inoculation were surface-sterilized and washed in approximately 0.01% Tween 20 (5 min), 0.25% sodium hypochlorite (2 min), 70% ethanol (30 sec), distilled water (4 x 10 sec). Leaf discs were cut from a leaf in pairs; one was placed on Murashige and Skoog (MS) medium alone and the other on kanamycin sulphate-supplemented MS medium. Plates were incubated at 32°C with a photoperiod of 16 hours. Leaf discs were transferred to freshly prepared medium every seven days.

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Mechanical inoculation of *N. benthamiana* plants with in vitro transcripts derived from DNA constructs pTB2, pTBD4 and pTBN62, respectively, resulted in symptomatic infection with virus of typical TMV shape and yield (1.5-5.8 mg virus/g tissue). Symptoms were less severe compared to TMV-UI-infected plants and consisted of plant stunting with mild chlorosis and distortion of systemic leaves. The sizes of virion RNA from systemically infected tissue of plants inoculated with TB2, TBD4 and TBN62, respectively, were consistent with predicted lengths of RNA transcribed in vitro from the respective plasmids. These RNA species contained TMV sequences plus their respective bacterial gene inserts. Probes complementary to the manipulated portion of the respective genomes were protected in RNase protection assays by progeny TBD4 and TBN62 viral RNAs. This indicated that the precise and rapid deletion of inserted sequences which had been a problem with previous constructs (11) did not occur with TBD4 or TBN62. It was hypothesized that with the previously reported constructs, foreign inserts were deleted due

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to recombination between repeated subgenomic promoter sequences (11) With TBD4 and TBN62, such repeated sequences were reduced by employing heterologous subgenomic mRNA promoters. Additional bands that were seen and were smaller than the probe and smaller than the full-length viral RNA might represent alterations within a portion of the TBN62 population, although in this case the relative proportion of full-length and additional smaller bands was unchanged following a subsequent passage.

The sequence stability of TBD4 and TBN62 virion RNA was examined in serial passages through N. benthamiana. Plants were inoculated with two and four independent in vitro transcription reactions from pTBD4 and pTBN62, respectively, and systemically infected leaf tissue was serially passaged every 11-12 days. After 48 days of systemic infection, full-length virion RNA of TBD4 including the DHFR sequences was still detected by Northern transfer hybridization, and still protected probes complementary to the manipulated portion of the genome in RNase protection assays. Five clonal populations of virion RNA were derived from TBD4-infected plants propagated for 170 days (one series involving 10 passages) by isolation of local lesions on N. tabacum Xanthi-nc. The consensus DHFR sequence for three of the populations corresponded with the published DHFR sequence except for a translationally silent third base change (U→C) at nucleotide 72 of the coding sequence. The nucleotide change at position 72 of the DHFR coding sequence was not evident in progeny RNA from TBD4 infected plants propagated for 48 days. Virion RNA from plants serially infected with TBN62 was less stable with different portions of the NPTII sequence being

14 deleted in each of the independent series of
43 passages. The time of loss of these sequences varied
between after the first passage (12-24 days) and the
third passage (36→47 days). The reason for the
occurrence of deletions in the NPTII sequence of
TBN62 is not known. However, on the basis of the
stability of the DHFR sequences in TB2, such
instability of inserted foreign sequences would not
seem to be an intrinsic feature of the expression
vector TB2. In contrast, such deletions might be
dictated by the nucleotide composition of the
inserted foreign sequences themselves. Similar
instabilities among DNA plant virus vectors have been
seen.

A commercial source of antiserum and sensitive
enzymatic assays for the extensively used selectable
marker NPTII (62) allowed further analysis of tissue
infected with TBN62. Western blot analysis, enzyme
activity, and leaf disc assays demonstrated the
presence of functional NPTII enzyme and its
phenotypic expression in plant tissue systemically
infected with TBN62 but not in TB2-infected or
healthy plants. NPTII protein and enzyme activity
was even detected in some TBN62-infected plants
propagated for 36 days.

It was evident that the levels of extractable
NPTII protein were considerably lower than coat
protein, the most highly expressed TMV protein. Such
low levels could be a reflection of the relative
stabilities or partitioning of the respective
proteins in plant cells, or might be due to one or
more aspects of the vector or foreign gene sequences
affecting the synthesis of subgenomic mRNA or post-
transcriptional expression of the reporter gene. The
relatively high yield of virus from plants infected

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with the vector constructs would seem to preclude a dramatic reduction in the efficiency of virus replication. However, one possibility for low expression might be the position of the reporter gene relative to the 3' terminus of the genome. The amount of the 30kDa protein produced by different mutants of TMV has been shown to be inversely proportional to the distance the 30kDa protein ORF was from the 3' terminus of the genome. This relationship was consistent with the observations of French and Ahlquist (63), i.e., that the level of subgenomic RNA from brome mosaic virus RNA 3 was progressively greater the closer the promoter was inserted to the 3' terminus.

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EXAMPLE 2

Although the RPM of Example 1 is capable of systemic spread in N. benthamiana, it is incapable of systemic spread in N. tabacum. This example describes the synthesis of RPM which is capable of systemic spread in N. tabacum.

The O-coat protein coding sequence contained in pTB2 was cut from pTB2 by digestion with AhaIII. The UI-coat protein coding sequence was removed from pTMV204 by digestion with AhaIII and inserted into AhaIII-digested pTB2 to produce vector pT8U5 (Fig. I)

The XhoI/SalI fragments from pNC4X (DHFR sequence) and pNU116 (NPTII sequence), respectively, are ligated into the XhoI site of pTBU5 in the same sense as the TMV coding sequences. N. tabacum plants are inoculated and analyzed as described in Example 1. Functional enzymes are seen in the systemically infected plants but not in the control plants.

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EXAMPLE 3

This example describes the synthesis of RPVNA in which the native coat protein gene is under control of its native subgenomic promoter and a non-native subgenomic promoter has been inserted to drive the expression of non-native nucleic acid.

The TMV-O promoter and the TMV-UI coat protein sequence are removed from pTB2 by digesting with XhoI and KpnI. The XhoI end is converted to a PstI site by blunt-ending and adding a PstI linker. This PstI/KpnI fragment is subcloned into a Bluescript vector. Two subclones of this Bluescript vector are created by site-directed mutagenesis as follows:

Bluescript Sub 1 is prepared by using PCT techniques to create a site-specific fragment that will force a mutation at the ATG (coat protein) start site and create a XhoI site near the ATG site.

Bluescript Sub 2 is prepared by using PCR techniques to create a site-specific fragment that will force a mutation at the TAA (coat protein) stop site and create a XhoI site near the TAA site. A PstI/XhoI cut of the Bluescript Sub 1 and a XhoI/KpnI cut of the Bluescript Sub 2 will give two fragments that can be ligated, giving a PstI/KpnI fragment that has a XhoI cloning insert site that is downstream from the TMV-O promoter. This PstI/KpnI fragment is inserted into the pTKUI vector that has had a NsiI/KpnI fragment removed. (PstI end can be ligated to NsiI). The resulting clone will be pTKU1-a with a TMV-O promoter on the 3' side and a XhoI insert site, into which can be inserted a gene-of-choice, that will be driven by the TMV-O promoter.

The XhoI/SalI fragments from pNC4X (DHFR sequence) and pNU116 (NPTII sequence), respectively, are ligated into the XhoI site of pTBU1-a in the same

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I sense as the TMV coding sequences. N. tabacum plants are inoculated and analyzed as described in Example 1. Functional enzymes are seen in the systemically infected plants but not in the control plants.

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EXAMPLE 4

Additional DNA coding sequences were prepared for insertion into RVPNAs having either the O-coat protein (Example 1) or the U1-coat protein gene (Example 2). In each instance, the coding sequence was synthesized to contain the XhoI site of pTB2 (Example 1) or pTBU5 (Example 2), in the same sense as the coding sequence.

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Standard procedures were used to transform the plasmids into E. coli and to isolate the DNA from an overnight culture. Following extraction of the plasmid DNA, an RNA copy of the TB2 or TBV5 vector (with or without the gene of choice) was made using a DNA-directed RNA polymerase. The RNA was capped during the reaction by adding m⁷GpppG₄ during the transcription reaction, as previously published. This RNA was then used to inoculate a tobacco plant. Standard virus isolation techniques can be used to purify large concentrations of the transient vector for inoculations of multiple numbers of plants.

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A coding sequence for Chinese cucumber α -trichosanthin containing XhoI linkers is shown in SEQ ID NO: 3, with the corresponding protein as SEQ ID NO: 4.

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A coding sequence for rice α -amylase containing XhoI linkers is shown in SEQ ID NO: 5, with the corresponding protein as SEQ ID NO: 6. This sequence was prepared as follows:

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The yeast expression vector pEno/I03 64 was digested with HindIII and treated with mung bean exonuclease to remove the single-stranded DNA overhang. The .16 kb HindIII (blunt end) fragment containing the entire rice α -amylase cDNA 05103 65 1990; GenBank accession number M24286) was digested with ScaI and linkerered with a XhoI oligonucleotide 40 (5' CCTCGAGG 3'). The modified α -amylase cDNA fragment was isolated using low-melt agarose gel electrophoresis, subcloned into an alkaline phosphatase treated XhoI site in pBluescript KS+(Stratagene, La Jolla, Calif.), and maintained in E. coli K-12 strain C-600.

A rice α -amylase coding sequence containing a short 3'-untranslated region was prepared as follows:

The E. coli vector pVC18/13 (64) was digested with KpnI, XhoI and treated with ExoIII and mung bean exonuclease. The modified plasmid was treated with DNA poll, DNA ligase, and transformed into C-600. An isolate, clone pUC18/3 #8, had a 3' deletion that was very close to the stop codon of 05103. This plasmid was digested with EcoRI, treated with mung bean exonuclease, and linkerered with a XhoI oligonucleotide 40 (5' CCTCGAGG 3'). A 1.4 Kb HindIII-XhoI fragment from 190 the resulting plasmid (pUC18/3 #8X) was isolated 190 using low melt agarose gel electrophoresis, subcloned into pBluescript KS- (Stratagene, La Jolla, Calif.) I and maintained in E. coli K-12 strains C-600 and JM109. The deletion was sequenced by dideoxy termination using single-stranded templates. The deletion was determined to reside 14 bp past the rice 190 α -amylase stop codon. Plasmid pUC18/3 #8X was digested with HindIII, treated with mung bean exonuclease, and linkerered with a XhoI oligonucleotide 40 (5 'CCTCGAGG 3') A 1.4 Kb XhoI fragment was isolated

by trough elution, subcloned into an alkaline phosphatase-treated XbaI site in pBluescript KS+, and maintained in JM109.

60, 62 A sequence containing the coding sequence for human α -hemoglobin or β -hemoglobin and transit peptide of petunia EFSP synthase is shown in SEQ ID NO: 7 or SEQ ID NO: 8, and corresponding protein sequences as SEQ ID NO: 9 and SEQ ID NO: 10, respectively.

60 **I** Purified protein extracts from N. benthamiana treated with a recombinant plant viral nucleic acid containing the gene for α -trichosanthin, prepared in accordance with Example 1, were separated using polyacrylamide gel electrophoresis and probed with antibodies specific for α -trichosanthin using standard procedures for Western analysis. Figure 2 is an autoradiograph of the gels which demonstrates production of processed α -trichosanthin protein in plants treated with a recombinant plant viral nucleic acid containing the gene for α -trichosanthin.

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EXAMPLE 5

Field Tests
The field site design contained two experiments (1 and 2). Experiment 1 was a typical row crop configuration that contained untreated border rows (8) of tobacco on all outside perimeter rows as well as internal rows. In addition, every fourth row was a spacer row (S) that was left unplanted in order to allow large farm equipment to access the field (e.g., for spraying pesticides) without coming into direct contact with any of the treated rows (T). Each inoculation was administered by direct hand application of the vector to a single leaf of an individual plant. No spray inoculum was used.

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Experiment 2 was a typical plantbed configuration. A high density of plants per square foot was grown at a uniform height by frequent clipping of the plantbed using a modified mower attached to a tractor power takeoff. This experiment contained a complete perimeter border of plantbeds that was not inoculated with the vectors. Inoculation of the treated plantbeds was made using a downward-directed spray through the modified mower blade assembly and administered so as to prevent overspray to adjacent plantbeds.

Experiment 1 was a split-plot design using row culture with seven genotypes as main plots in randomized blocks and four replications. Each plot was 13 feet long and consisted of three rows, with only the middle three or four plants of each center row used for testing. Rows were four feet on center and plants spaced 20 to 22 inches in the row.

Experiment 2 was a randomized complete block design using plantbed culture with four genotypes and three replications. Each plot consisted of a 4-foot by 12-foot plantbed.

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Genotypes. Experiment 1: (Nicotiana tabacum) K-326, Sp G-28, TI-560, Md-609, Galpao, Wisc-503B and Nicotiana benthamiana.

Experiment 2: (Nicotiana tabacum) K-326, TI-560, Md-609, Galpao.

Chemical Fertilization. Experiment 1: 800 lbs 6-12-18 after transplanting; 100 lbs 33-0-0 after first harvest; 200 lbs 15-0-14 after second harvest.

Experiment 2: 2400 lbs 12-6-6 at time of plantbed formation; 300 lbs 33-0-0 after first harvest; 670 lbs 15-0-14 after second harvest.

Clipping. Experiment 2 was clipped twice a week for two weeks, to impart uniformity to the plants.

Weed, Insect and Disease Control. Experiment 1:
Prior to forming rows, Paarlan 6B (1 qt/A), Temik 15G
(20lb/A) and Ridomil (2 qts/A) were broadcast-applied
and incorporated by disking. During row formation,
Telone C-17 (10.5 gal/A) was applied. After
178 transplanting, Dipel (1/2 lb/A) was applied to
170 control budworms and hornworms. Orthene (2/3 lb/A)
was applied to control aphids and hornworms as
necessary.

Experiment 2: Ridomil 2G (1 qt/A; 1 oz/150 sq
yds) was applied at seeding and at weekly intervals
14 beginning 60-70 days after seeding (as needed).
Carbamate 76WP (3 lb/100 gal water) was also used as
foliar spray as needed in the initial plantbed stage,
to control Anthracnose and Damping-off diseases. At
178 normal transplanting size, Dipel (1/2 lb/A) was
170 applied. Orthene (2/3 lb/A) was applied to control
aphids and hornworms as necessary.

Transplanting. Experiment 1 was transplanted
using seedlings pulled from the plantbeds of
Experiment 2.

Inoculation. Experiment 1: A single leaf on
each non-control plant was hand-inoculated with a
selected recombinant plant viral nucleic acid
60 containing NPT II, α -trichosanthin or rice α -amylase.
Each individual plant was inoculated with a single
vector.

Experiment 2: The plants were inoculated with
the vectors described in Experiment 1, using a spray
applied through the deck of the clipping mower while
the plants are being clipped a final time. Each non-
control plot received only a single vector construct.
Control plants received no inoculation with any
vector.

Data Collection. Experiment 1: Sampling of both inoculated and control plant leaves was conducted on a schedule (approximately weekly) during first growth until plants were approximately 30 inches tall. Plants were then cut (harvest 1) with a rotary brush blade to leave six inches of stalk exposed above the ground. The plants were then allowed to continue growth (second growth) to a height of approximately 30 inches. Leaf samples were taken just before harvest 2. This procedure for cutting, growth and sampling was repeated for third growth and for fourth growth, if detectable amounts of the genes of interest inserted into the vectors were found.

Experiment 2: Sampling of 10 plants from each plot was conducted on a schedule (approximately weekly) from inoculation to harvest 1 and from harvest 1 until harvest 2. Following harvest 2, sampling was conducted only at harvest 3.

Sample Size and Analytical Methods. A 1.6 cm disk was excised from a single leaf near the apex of the plant. Each leaf disk was placed either in a 25 ml glass vial with screw cap and containing absolute ethanol or in a sealable plastic bag.

Leaf discs were either preserved in absolute ethanol or lyophilized. Depending on the specific gene product to be detected, leaf samples were prepared according to standard techniques for Northern or Western blot analyses or specific enzyme activity.

During first growth, visual monitoring of the plants treated with the RPVNA were conducted to observe any external phenotypic expression of the vector system. In some cases, the phenotypic expression was typical of Tobacco Mosaic Virus infections (lighter

and darker "mosaic" patterns in the leaf). In other cases, the only symptoms seen were on the inoculated leaf, which included white or brown speckels of approximately 2mm in diameter and/or suppression of the central vein elongation of the leaf.

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EXAMPLE 6

P A full-length DNA copy of the OMV genome is prepared as described by Dawson, W.O. et al. (4). The vector containing the DNA copy of the OMV genome is digested with the appropriate restriction enzymes or suitable exonucleases to delete the coat protein coding sequence. The deletion of the coding sequence for the viral coat protein is confirmed by isolating OMV and using it to infect germinating barley plants. The isolated OMV RNA is incapable of spreading beyond the lesion under natural conditions. A vector containing the OMV sequences is prepared as described in Examples 1-3.

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EXAMPLE 7

P A full-length DNA copy of the genome is prepared as described by Dawson, W.O. et al. (4). The vector containing the DNA copy of the ENV genome is digested with the appropriate restriction enzymes or suitable exonucleases so as to delete the coat protein coding sequence. The deletion of the coding sequence for the viral coat protein is confirmed by isolating RNV RNA and using it to infect germinating barley plants. The isolated is incapable of spreading beyond the lesion under natural conditions. A vector containing the OMV sequences is prepared as described in Examples 1-3.

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EXAMPLE 8

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A full-length DNA copy of the PVY or PVX genome is prepared as described by Dawson, W.O. et al. (4). The vector containing the DNA copy of the PVY or PVX genome is digested with the appropriate restriction enzymes or suitable exonucleases to delete the coat protein coding sequence. The deletion of the coding sequence for the viral coat protein is confirmed by isolating PVY or PVX ENA and using it to infect potato plants. The isolated PVY or PVX RNA is incapable of spreading beyond the lesion under natural conditions. A vector containing the OMV sequences is prepared as described in Examples 1-3.

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EXAMPLE 9

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A full-length DNA copy of the maize streak virus (MSV) genome is prepared as described by Dawson, W.O. et al. (4). The vector containing the DNA copy of the MsV genome is digested with appropriate restriction enzymes or suitable exonucleases to delete the coat protein coding sequence. Deletion of the coding sequence for the viral coat protein is confirmed by isolating MSV and using it to infect potato plants. The isolated MSV is incapable of spreading beyond the lesion under natural conditions. A vector containing the OMV sequences is prepared as described in Examples 1-3.

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EXAMPLE 10

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A full-length DNA copy of the TGMV genome is prepared as described by Dawson, W.O. et al. (4).

The vector containing the DNA copy of the TGMV genome is digested with the appropriate restriction enzymes or suitable exonucleases to delete the coat protein coding sequence. The deletion of the coding sequence for the viral coat protein is confirmed by isolating TGMV RNA and using it to infect potato plants. The isolated TGMV RNA is incapable of spreading beyond the lesion under natural conditions. A vector containing the TGMA sequences is prepared as described in Examples 1-3.

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EXAMPLE 11

The coding sequence for beta-cyclodextrin glucotransferase is isolated from alkalophilic Bacillus sp. strain No. 38-2 in the following manner:

The chromosomal DNA of strain No. 38-2 (66) is partially cleaved with Sau3AI, and the fragments ligated in BamHI-digested pBR322. A transformant carrying plasmid pCS115, which contains a 3.2 kb DNA fragment from the genome of the producing strain, has the CGT activity. The CGT produced by this transformant gives one line of precipitation which fuses completely with that for the No. 38-2 CGT by an Ouchterlony double-diffusion test. The nucleotide sequence of the fragment is found by the dideoxy chain termination reaction using pUC19, and the exonuclease deletion method (67). The nucleotide sequence of the fragment shows a single open reading frame corresponding to the CGT gene. A protein with a molecular mass of 66 kDa could be translated from this open reading frame of 1758 bp. For the detailed nucleotide sequence, see Hanamoto, T. et al. (66).

The sequence of the N-terminal amino acids of the extracellular form of CGT is found with a peptide

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sequencer. NH₂-Ala-Pro-Asp-Thr-Ser-Val-Ser-A5n-Lys-Gln-Asn-Phe-Ser-Thr-Asp-Val-Ile (SEQ ID NO: 6) is identical to that deduced from the DNA sequence (residues 1 to 17). This result suggests that 27 amino acid residues (residues -27 to -1) represent a signal peptide which is removed during secretion of CGT. The molecular weight of the matured CGT calculated from the DNA sequence is 63,318.

A probe is prepared based on a portion of the amino acid sequence of cyclodextrin glucanotransferase and used to isolate the coding sequence for this enzyme. Alternatively, the beta cyclodextrin glucotransferase coding sequence is isolated following reverse transcription. The fragment containing the coding sequence is isolated and cloned adjacent the subgenomic promoter of the native viral coat protein gene in the vectors prepared in Examples 6-10.

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EXAMPLE 12

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The RPVNA of Example 11 is used to infect corn plants (viruses based on OMV, RNV, or TGMV) or potato plants (viruses based on PVY or PVX). The infected plants are grown under normal growth conditions. The plants produce cyclodextrin glucotransferase which catalyzes the conversion of starch to cyclodextrin in the plant tissue. The cyclodextrin is isolated by conventional techniques.

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EXAMPLE 13

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A. The coding sequence for an esterase is isolated from Bacillus subtilis Thai 1-8 (CBS 679.85) as follows. The positive selection vector pUN121

(68) is used. This vector carries an ampicillin resistance gene, a tetracycline resistance gene and a C₁-repressor gene. Transcription of the tetracycline gene is prevented by the gene product of the C₁-repressor gene. Insertion of foreign DNA into the H BclI site of the C₁-repressor gene results in activation of the tetracycline gene. This allows positive selection of recombinants on ampicillin/tetracycline agar plates.

I Partially Sau3a-digested Bacillus subtilis Thai 1-8 DNA is mixed with BclI-digested pUN121 DNA.

I After recirculation by the use of polynucleotide ligase, the DNA mixture is introduced into E. coli DH1 (ATCC No. 33849) using the CaCl₂ transformation procedure.

I One thousand E. coli colonies are obtained which are resistant to ampicillin and tetracycline. All transformants are stored and replica-plated according to Gergan et al. (69). Replicated colonies are screened using a soft agar overlay technique, based on a previously described procedure to detect esterase activity (70).

62 Essentially, a mixture of 0.5% low-melting agarose, 0.5M potassium phosphate (pH 7.5), 0.5 mg/l β-naphthyl acetate and 0.5 mg/ml fast-blue is spread over the transformants. Within a few minutes, colonies with esterase or lipase activity develop purple color. Such colonies are grown overnight in

H 2^x YT (16 g/l Bactotryptone, 10 g/l yeast extract, 5 g/l NaCl) medium and subsequently assayed for their ability to convert S-naproxen ester to S-naproxen

T (the method of Example 1 of EP-A 0233656). One E. coli transformant is able to convert S-naproxen ester. The plasmid isolated from this transformant, which is called pNAPT-2 (CBS 67186). Its size is 9.4 kb.

HindIII restriction enzyme fragments of pNAPT-2 are ligated into pPNEO/ori. This is performed as described below. pPNEO/ori is constructed by ligating the 2.7 kb EcoRI/SmaI restriction fragment of pUC19 to the 2.5 kb EcoRI-SnaBI restriction fragment of pUB110. The resulting shuttle plasmid, pPNEO/ori (5.2 kb) has the capacity to replicate both in E. coli and in Bacillus species due to the presence of the pUC19 origin, and the pUB110 origin. In addition, pPNEO/ori carries a gene encoding ampicillin resistance and a gene encoding neomycin resistance.

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For subcloning, HindIII-digested pNAPT-2 is mixed with HindIII-digested pPNEO/ori and ligated. The mixture is transformed to E. coli JM101 hsds as described (Maniatis et al., supra). E. coli JM101 hsds is obtained from the Phabagen collection (Accession No. PC 2493, Utrecht, The Netherlands). Colonies capable of hydrolyzing β -naphthyl acetate are selected as described in Example 56 of EPA 0 233 656. From two positive colonies, pNAPT-7 and pNAPT-8 plasmid DNA is isolated and characterized in detail by determining several restriction enzyme recognition positions.

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B. The coding sequence for an E. coli esterase is prepared as follows:

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Plasmids pIP1100 (isolated from E. coli BM 2195) and pBR322 are mixed, digested with AvaI, ligated and transformed into E. coli, and clones are selected on Em (200 /g/ml). Transformants resistant to Ap and Em but also to Sm are analyzed by agarose gel electrophoresis of crude lysates. The transformant harboring the smallest hybrid plasmid is selected, its plasmid DNA is digested with AvaI, and the 3.5 kb pIP1100 insert is purified and partially digested

with Sau3A. The restriction fragments obtained are cloned into the BamHI site of pBR322 and transformants selected on Em are replica-plated on Sm. The plasmid content of transformants resistant only to Ap and Em is analyzed by agarose gel electrophoresis. DNA from the smallest hybrid, pAT63, is purified and analyzed by agarose gel electrophoresis after digestions with Sau3A, EcoRI, PstI or HindIII-BamHI endonucleases (not shown). Plasmid pAT63 consists of pBR322 plus a 1.66 kb pIP1100 DNA insert. Purified EcoRI-HindIII (1750-bp) and BamHI-PstI (970-bp) fragments of pAT63 are subcloned into pUC8 and found not to confer resistance to Em.

The HpaII-BamHI fragment of pAT63 is sequenced according by the Sanger technique. The complete sequence is shown in Ounissi, H. et al. (71).

C. The coding sequence from acylase is isolated from Arthrobacter viscosus 8895GU, ATCC 27277 follows.

A gene library of A. viscosus 8895GU is constructed by inserting EcoRI-cleaved A. viscosus chromosomal DNA into the EcoRI cleavage site of pACYC184. The vector DNA and A. viscosus DNA are both digested with EcoRI. The 5' end of the vector DNA is dephosphorylated with calf intestinal alkaline phosphatase. Deposphorylated vector DNA and digested A. viscosus DNA are incubated with T4 DNA ligase and transformed into E. coli HB101.

Transformed colonies of E. coli were screened by the Serratia marcescens overlay technique. Penicillin G was added to the medium. S. marcescens is sensitive to the deacylation product of penicillin G, 6-aminopenicillamic acid (6-APA). Colonies of transformed E. coli will produce areas of S.

I L marcescens inhibition in overnight cultures. The plasmid carried by transformed E. coli is referred to as pHYM-1. The plasmid having opposite DNA orientation is designated pHYM-2 (72).

D. A coding sequence for human gastric lipase mRNA is prepared by guanidinium isothiocyanate extraction of frozen tissue. Polyadenylated RNA is isolated by oligo(dT)-cellulose chromatography. cDNA is prepared from human stomach mRNA by procedures well known in the art. cDNA is annealed to PstI-cut dG-tailed pBR322. The hybrid plasmid is transformed into E. coli DH1. Transformants are screened by colony hybridization on nitrocellulose filters. The probe used is synthesized from the rat lingual lipase gene and labeled by nick translation. Positive colonies are grown up and plasmids are analyzed by restriction endonuclease mapping.

14 An esterase acylase or lopase gene prepared as described above is removed from the appropriate vector, blunt-ended using mung bean nuclease or DNA polymerase I, and XhoI linkers added. This esterase with XhoI linkers is cleaved with XhoI and inserted into the vertors described in Examples 1-3 or 6-10. Infection of the appropriate host plants by the RPVNA prepared in accordance with Example 2 results in the synthesis of esterase, acylase or lipase in the plant tissue. The enzyme is isolated and purified by conventional techniques and used to prepare stereo-specific compounds. O

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EXAMPLE 14

The coding sequence for CMS-T is isolated from a BamHI maize mtDNA library as described by Dewey, R.E., et al. (73). The ORF-13 coding sequence is

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40 isolated by restriction endonuclease digestion followed by 5'-exonuclease digestion to the start codon. alternatively, a restriction site is engineered adjacent the start codon of the ORF-13 coding sequence by site-directed oligonucleotide mutagenesis. Digestion with the appropriate restriction enzyme yields the coding sequence for ORF-13. The fragment containing the ORF-13 coding sequence is isolated and cloned adjacent the promoter of the native viral coat protein gene in the vectors prepared in Examples 6, 7 and 10.

Maize plants are infected by teh RPVNA prepared in accordance with Example 1. The infected plants are grown under normal growth conditions. The plants produce cms-T which induces male sterility in the infected maize plants.

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EXAMPLE 15

P H The coding sequence of S₂-protein (for self-
I incompatibility) is isolated from Nicotiana alata as
H described in EP-A 0 222 526. The S₂-protein coding
sequence is isolated by restriction endonuclease
40 digestion followed by 5'-exonuclease digestion to the
start codon. Alternatively, a restriction site is
H engineered adjacent the start codon of the S₂-protein
coding sequence by site-directed oligonucleotide
mutagenesis. Digestion with the appropriate
restriction enzyme yields the coding sequence for
H S₂-protein. The fragment containing the S₂-protein
coding sequence is isolated and cloned adjacent the
14 promoter of the viral coat protein gene in the
vectors prepared in Examples 1-3.

Tobacco plants are infected by the RPVNA prepared in accordance with Example 1, prior to pollen formation. The infected plants are grown under normal growth conditions. The plants produce S-protein which induces male sterility via the self-incompatibility mechanism.

The following example demonstrates that high levels of therapeutic proteins can be expressed using the plant RNA viral vectors of the present invention.

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EXAMPLE 16

Rapid and High Level Expression of Biologically Active α -trichosanthin in Transfected Plants Using a Novel RNA Viral Vector

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Trichosanthin is a eukaryotic ribosome inactivating protein found in the roots of a Chinese medicinal plant (74). In Trichosanthes kirilowii

I 60 Maximowicz, α -trichosanthin is a monomeric protein which catalyzes the cleavage of an N-glycosidic bond in 28S rRNA (75,76). This reaction inhibits protein synthesis by affecting the ability of the 60S ribosomal subunit to interact with elongation factors. The mature compound has an approximate relative molecular mass of 27 kDa and is initially produced as a preprotein (77). During its biosynthesis, a putative 23 amino acid secretory signal peptide is removed and a 19 amino acid peptide is probably excised from the carboxy terminus.

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Purified T. kirilowii derived α -trichosanthin causes a concentration-dependent inhibition of HIV replication in acutely infected CD4+ lymphoid cells, and in chronically infected macrophages (78,79). This compound is currently being evaluated in clinical studies as a potential therapeutic drug in

the treatment for HIV infection (80). The exact
60 mechanism of anti-HIV infection by α -trichosanthin is
not known. Amino acids involved in catalysis and
inhibition of HIV replication may be identified using
site directed mutagenesis. Detailed
structure/function analysis will require an abundant
source of recombinant protein as well as a rapid
method for generating and analyzing mutants.

60I Although the expression of α -trichosanthin in E. coli
has been reported previously (81, 97), the amount
synthesized was low (approximately 0.01% total
cellular protein), the carboxy terminal extension was
not removed, and the biological activity of the
compound was not determined.

Tobamoviruses, whose genomes consist of one
plus-sense RNA strand of approximately 6.4 kb, have
been used to produce heterologous proteins. RNA
transcripts from viral cDNA clones serve as
infectious templates, encoding proteins involved in
RNA replication, movement, and encapsidation (82).
Subgenomic RNA for messenger RNA synthesis is
controlled by internal promoters located on the
minus-sense RNA strand (83). TMV RNA viruses have
been used previously to express Leu-enkephlin in
tobacco protoplasts (84) and bacterial
chloramphenicol acetyltransferase in inoculated
tobacco leaves (85,86). These previous attempts to
express foreign genes have resulted in either
unstable constructs or loss of long distance viral
movement. Recently, Nicotiana benthamiana plants
transfected with a hybrid virus consisting of tobacco
mosaic virus, strain U1 (TMV-U1) and an additional
RNA subgenomic promoter from odontoglossum ringspot
virus (ORSV) produce a systemic and stable expression
of neomycin phosphotransferase (87).

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Construction of pBGC152

The plasmid pSP6-TKU1 contains the entire TMV-U1 genome fused to the SP6 promoter by oligonucleotide directed mutagenesis and inserted into pUC118 as a XhoI/KpnI fragment. The sequence of the mutagenesis primer used to attach the SP6 promoter sequence to the TMV genome is:

40 L 5' -GGGCTCGAGATTTAGGTGACACTATAGTATTTACAACATTACCA-

3' wherein the XhoI site is in italics, the SP6 promoter is in boldface and the TMV sequence is underlined. The primer was attached to a TMV subclone I14 called pC48 (Raffo, et al., Virology 184: 277-289 (1991)). The promoter was attached by PCR using the above primer and a primer complementary to TMV sequences 5673 to 5692. This amplification produced a fragment of ca. 614bp, which was then digested with XhoI and EcoRI (TMV 270) to produce a ca. 292 bp fragment which was then subcloned into similarly cut pUC129 resulting in plasmid pSP6-T1.

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pSP6-T1 was cut with XhoI and XmaI (a SmaI isoschizomer which cuts at TMV 256) and the resulting ca. 278 bp fragment was ligated into pTKU1 (Donson, et al. Proc. Natl. Acad. Sci. U.S.A. 88:7204-7208 (1991)) which had been modified by cutting at the unique PstI site at the 5' end of the genome, blunting with T4 DNA polymerase, followed by the addition of XhoI linkers. This resulted in the infectious clone pSP6-TKU1 and XmaI digested.

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As shown in FIG. 7, the EcoRI site in pBR322 was mutagenized to a KpnI site using EcoRI, DNA polymerase (Klenow), and KpnI linkers. A KpnI\BamHI fragment of the resulting plasmid, pBSG121, was substituted with a KpnI\BamHI fragment of pTB2 (ATCC No. 75,280 deposited July 24, 1992). A SalI/KpnI fragment of the resulting plasmid, pBSG122, was

substituted with a XhoI/KpnI fragment of pSP6-TKUI (also known as T1) which resulted in plasmid pBGC150.

A BamHI/KpnI fragment of pBGC150 was substituted with a BamHI/-KpnI fragment of pTB2/Q resulting in plasmid pBGC152. pTB2/Q was constructed beginning with plasmid pQ21D (ATCC No. 67907) described in F Piatak, Jr., et al. U.S. Patent No. 5,128,460, the contents of which are incorporated herein by reference. The plasmid "clone 5B" containing a PCR amplified 0.88 kb XhoI fragment of the TCS sequence in pQ21D was obtained using oligonucleotide mutagenesis to introduce XhoI cloning sites at the start and stop codons of pQ21D such that the following sequence was obtained: 5'-CTCGAGGATG ATC - 40 L --- // --- ATT TAG TAA CTCGAG-3' (XhoI site in italics). A 0.88 kb XhoI fragment from "clone B" was subcloned into the XhoI site of plasmid pTB2 in the sense orientation to create plasmid pTB2/Q. C

CL In vitro transcriptions, inoculations,
and analysis of transfected plants

PI N. benthamiana plants were inoculated with in vitro transcripts of KpnI digested pBGC152 as described previously (89). Virions were isolated I from N. benthamiana leaves infected with BGC152 transcripts, stained with 2% aqueous uranyl acetate, and transmission electron micrographs were taken using a Zeiss CEM902 instrument.

CL 60 Purification, immunological detection,
and in vitro assay of α -trichosanthin

P Two weeks after inoculation, total soluble protein was isolated from 3.0 grams of upper, non- I inoculated N. benthamiana leaf tissue. The leaves were frozen in liquid nitrogen and ground in 3 mls of 5% 2-mercaptoethanol, 10 mM EDTA, 50 mM potassium

phosphate, pH 6.0. The suspension was centrifuged
60 and the supernatant, containing recombinant α -
trichosanthin, was loaded on to a Sephadex G-50
column equilibrated with 2 mM NaCl, 50 mM potassium
phosphate, pH 6.0. The sample was then bound to a
Sephadex-S Fast Flow ion exchange column. Alpha-
trichosanthin was eluted with a linear gradient of
14 0.002-1 M NaCl in 50 mM potassium phosphate, pH 6.0.
60 Fractions containing α -trichosanthin were
concentrated with a Centricon-20 (Amicon) and the
buffer was exchanged by diafiltration (Centricon-10,
50 mM potassium phosphate, pH 6.0, 1.7 M ammonium
sulfate). The sample was then loaded on a HR5/5
14 alkyl superose FPLC column (Pharmacia) and eluted
with a linear ammonium sulfate gradient (1.7-0 M
ammonium sulfate in 50 mM potassium phosphate, pH
6.0). Total soluble plant protein concentrations
were determined (90) using BSA as a standard. The
60 concentration of α -trichosanthin was determined using
H the molar extinction coefficient of $E_{280} = 1.43$. The
purified proteins were analyzed on a 0.1 % SDS, 12.5%
polyacrylamide gel (91) and transferred by
electroblotting for 1 hour to a nitrocellulose
membrane (92). The blotted membrane was incubated
60 for 1 hour with a 2000-fold dilution of goat anti- α -
trichosanthin antiserum. The enhanced
chemiluminescence horseradish peroxidase-linked,
3 rabbit anti-goat IgG (Cappel) was developed according
to the manufacturer's (Amersham) specifications. The
21 autoradiogram was exposed for <1 second. The
60 quantity of total recombinant α -trichosanthin in an
extracted leaf sample was determined by comparing the
crude extract autoradiogram signal to the signal
obtained from known quantities of purified GLQ223.
The ribosome inactivating activity was determined by

measuring the inhibition of protein synthesis in a rabbit reticulocyte lysate system.

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Confirmation of High Level Expression
of Biologically Active α -trichosanthin

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The plant viral vector of the present invention
60 directs the expression of α -trichosanthin in
transfected plants. The open reading frame (ORF) for
60 α -trichosanthin, from the genomic clone pQ21D (88),
was placed under the control of the tobacco mosaic
virus (TMV) coat protein subgenomic promoter.

Infectious RNA from pBGC 152 (Fig. 3) was prepared by
in vitro transcription using SP6 DNA-dependent RNA
polymerase and were used to mechanically inoculate
I N. benthamiana. The hybrid virus spread throughout
all the non-inoculated upper leaves as verified by
transmission electron microscopy (Fig. 4), local
lesion infectivity assay, and polymerase chain
reaction (PCR) amplification (20; data not shown).

60 The 27 kDa α -trichosanthin accumulated in upper
leaves (14 days post inoculation) to levels of at
least 2% of total soluble protein and was analyzed by
immunoblotting, using GLQ223 (78), a purified

I 60 T. kirilowii derived α -trichosanthin, as a standard
(Fig. 5A). No detectable cross-reacting protein was

I observed in the non-infected N. benthamiana control
plant extracts (Fig. 5A, lane 5). Recombinant
60, 62 α -trichosanthin was easily detected in 7 μ g of crude
leaf extract using a Coomassie stain (Fig. 5B,
lane 3).

Prior investigators have reported a maximum
accumulation of a foreign protein in any genetically
engineered plant of 2% of the total soluble protein.
Although the expression of potentially valuable
proteins such as antibodies and human serum albumin
has been reported previously (94,95) these were

produced in Agrobacterium-mediated transgenic plants. A major difference between this plant viral expression system and previous methods is the quantity of protein produced and the amount of time required to obtain genetically engineered plants.

60 Systemic infection and expression of α -trichosanthin occurred in less than two weeks while it takes several months to create a single transgenic plant.

60 1 The α -trichosanthin produced and purified from upper leaves in transfected N. benthamiana (14 days post inoculation) was structurally identical to

60 L native α -trichosanthin. The 27 kDa protein cross-reacted with anti- α -trichosanthin antibody and had an identical FPLC purification profile as the GLQ223 standard. Although the C-terminal sequence of the recombinant protein was not analyzed, both GLQ223 and the purified recombinant α -trichosanthin appeared to have identical electrophoretic mobilities (Fig. 5B).

60 The exact C-terminal amino acid of the recombinant α -trichosanthin remains to be determined. The N-terminal sequence, Asp-Val-Ser-Phe-Arg-Leu-Ser was obtained from the purified protein using an automated protein sequenator (96). This result indicated that the putative signal peptide of the preparation was correctly processed at the site indicated in Fig. 1. The removal of the putative signal peptide at this site was consistent with the statistical expectation by the method of von Heijne (97). It is possible

60 that the α -trichosanthin signal peptide contributed to its high level expression by targeting the protein into the extracellular space. The nucleotide

60 sequences surrounding the α -trichosanthin start codon might also have an effect on the efficiency of translation initiation.

It is interesting to note that nucleotides flanking the translation initiating sites of the
40 highly expressed TMV-U1 (5' TTAAATATGTCT 3') and ORSV
L (5' TGAAATATGTCT 3') coat protein genes are conserved
40 while the corresponding region in pBGC152/α-
40 trichosanthin (5' TCGAGGATGATC 3') shows very little
similarity. It is possible that site directed
mutagenesis of nucleotides near the translation
60 initiation site of α-trichosanthin might increase its
expression.

60 The recombinant α-trichosanthin caused a
concentration dependent inhibition of protein
synthesis in a cell-free rabbit reticulocyte
H translation assay (Fig. 6). The ID₅₀ (dosage required
10 for 50% inhibition) was approximately .1 ng/ml, a
I value comparable to T.kirilowii derived
60 α-trichosanthin (GLQ223). Based on the ID₅₀ and dose
response, the enzyme produced in transfected plants
had the same specific activity as the native protein.
This result suggests that the fidelity of the viral
RNA-dependent RNA polymerase was relatively high
since base pair substitutions and deletions in the
foreign sequence during viral amplification would
lower the specific activity of the recombinant
enzyme.

As the disclosed and claimed invention
demonstrates, pBGC152 can direct the heterologous
60 expression of biologically active α-trichosanthin in
transfected plants. Large scale production of
recombinant proteins can be easily obtained using the
RNA viral-based system by simply increasing the size
and number of inoculated plants. Since tissue
60 containing high concentrations of α-trichosanthin can
be harvested two weeks after inoculation this system
can be used to rapidly screen the effects of site

(60) directed mutations. Identification of important amino acids involved in the inhibition of HIV replication in vivo may help to improve the efficacy of α -trichosanthin as a potential AIDS therapeutic drug.

The following plasmids have been deposited at the American Type Culture Collection (ATCC), Rockville, MD, USA, under the terms of the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedure and Regulations thereunder (Budapest Treaty) and are thus maintained and made available according to the terms of the Budapest Treaty. Availability of such plasmids is not to be construed as a license to practice the invention in contravention of the rights granted under the authority of any government in accordance with its patent laws.

The deposited cultures have been assigned the indicated ATCC deposit numbers:

	<u>Plasmid</u>	<u>ATCC No.</u>
T,870X a P	pTB2	75280
	pTB45	75281

While the invention has been disclosed in this patent application by reference to the details of preferred embodiments of the invention, it is to be understood that this disclosure is intended in an illustrative rather than limiting sense, as it is contemplated that modifications will readily occur to those skilled in the art, within the spirit of the invention and the scope of the appended claims.

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